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GenCore version 5.1.6
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July 29, 2004, 13:23:21; Search time 52 Seconds (without alignments) 168.442 Million cell updates/sec Run on:

US-09-852-238A-5

Title: Perfect score:

164 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR Sequence:

31

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

1586107 seqs, 282547505 residues Searched:

1586107 Total number of hits satisfying chosen parameters:

length: 0 length: 2000000000 Minimum DB seq Maximum DB seq

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

geneseqp2003as:\* geneseqp2003bs:\* geneseqp2004s:\* geneseqp1980s:\* geneseqp1990s:\* geneseqp2000s:\* A\_Geneseq\_29Jan04:\* geneseqp2001s:\* geneseqp2002s:\* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## STIMMARTES

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SUMMARIES	Ħ	AAW39910	AAW88231	AAW27406	AAW27408	AAW88238	ADC10142	ADC10144	AAW27407	AAW27123	AAW23835	AAW88232	AAE07048	AAG80111	AAE04321	AAE07039	AAB46858	ABB56342	AAB83354	AAB82948	AAU97152	AAM52829	AAM52828	ABG70597	ABG92883	AAE25811
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G-prot Human	Abr58602 Human can Aao29514 Human Can	Human	Human	Human	Aay41280 Fusion pr Aag80055 Chemokine		Abg32539 Human CCR Abg32540 Human CCR	Alanin	Aaw46973 Alanine s	Aaw46974 Alanine s	Aaw07602 Human G-p	Aay80128 Human G-p
ABB81054 ABB08343 ABC75540	ABR58602 AAO29514	ABU61654 ABP97728	ABP81933 ADC03341	AAW23834	AAY41280 AAG80055	AAW46976	ABG32539 ABG32540	AAW46975	AAW46973	AAW46974	AAW07602	AAY80128
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25 27 28	30	31 32	3 3 4	35	37	80 6	40	41	42	43	44	45

## ALIGNMENTS

Chemokine receptor; C-C CKR-5; Human Immunodeficiency Virus; HIV-1; CD4+ cell; inhibition; HIV-1 infection; beta-chemokine; non-synctium-inducing HIV-1 strain; treatment. Peptide representing an extracellular domain of C-C CKR-5. AAW39910 standard; peptide; 31 AA. (first entry) 06-JUL-1998 AAW39910; RESULT 1 

Homo sapiens. Synthetic

WO9747319-A1. 18-DEC-1997.

(AARO-) AARON DIAMOND AIDS RES CENT. 96US-0019941P. 96US-00665090. 97WO-US010619. (PROG-) PROGENICS PHARM INC 13-JUN-1997; 14-JUN-1996; 14-JUN-1996;

Moore JP, Litwin VM, Maddon PJ, Dragic T, WPI; 1998-086551/08. Allaway GP,

Trkola A;

Chemokine receptor CCR5 fragments - useful for inhibition of Human Immunodeficiency Virus 1 infection.

Disclosure; Page 2; 106pp; English.

Synthetic peptides AAW39910-13 represent the 4 extracellular domains of thuman chemokine receptor C-C CKR-5. C-C CKR-5 is capable of inhibiting the fusion of Ruman Immunodeficiency Virus (HIV)-1 to CD4+ cells and thus inhibiting HIV-1 infection of the cells. It is suggested that C-C CKR-5 functions as a beta-chemokine-sensitive second receptor for primary, non-synctium-inducing HIV-1 strains. The synthetic peptides were tested for their ability to inhibit membrane fusion mediated by the envelope glycoproteins of the LAI or JR-FL strains of HIV-1 using a resonance energy transfer assay. Specific inhibition of fusdion mediated by the JR- EL envelope glycoproteien was seen using the ECL2 peptide but not other

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Gaps

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Indels

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Length 100;

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The present sequence is an inactive human (C (Cys-Cys) chemokine receptor 5 (CCRS), which is not a receptor of human immunodeficiency virus type 1 or type 2 (CRS or its CDNA can used to diagnose, treat and/or prevent inflammatory diseases, e.g. rheumatoid arthritis, glomerulonephritis, asthma, idiopathic pulmonary fibrosis and psoriasis, viral infections, especially HIV-1 or HIV-2 infection, cancer, atherosclerosis and autoimmune disorders. Subjects that express the inactive receptor have a predisposition, or resistance to HIV-1 and/or
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Inactive; human Cys-Cys chemokine receptor-5; CCR5; human immunodeficiency virus; type 1; type 2; HIV-1; HIV-2; predisposition; resistance; diagnosis; treatment; prevention; inflammatory disease; rheumatorid arthritis; glomerulonephritis; asthma; idiopathic pulmonary fibrosis; psoriasis; viral infection; cancer; atherosclerosis; autoimmune disorder.
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100.0%; Pred. No. 7.9e-17;
ive 0; Mismatches 0;
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0; Mismatches 0;
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N-PSDB; AAT90116.
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Matches 31;
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CCRS (see AAW88232), but lacks transmembrane domains of wild-type CCRS (see AAW88232), but lacks transmembrane domains of wild-type CCRS (see AAW88232), but lacks transmembrane domains of strains of a co-receptor for infection by macrophage-tropic (M-tropic) strains of the wild type CCRS allele in an individual shows a positive correlation with resistance to infection with M-tropic HIV-1 strains, and may with resistance to infection with M-tropic HIV-1 strains, and may with resistance to infection with M-tropic HIV-1 strains, and may with a slower progression of the disease. The disease. The detection of CCRS variants may be used to identify individuals at lower risk of infection calative to the general population who, if infected, may exhibit slower progression to AIDS. Probes and primers (see AAV84127-36) are provided for use in diagnostic methods for detecting the presence of such variants. A method is provided for inhibiting HIV-1 infection of a cell expressing the CCRS receptor. This involves introducing a nucleic acid encoding a CCRS walled to the cell, thereby reducing the number of functional CCRS molecules present on the cell surface
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peptides. Polypeptides capable of inhibiting the fusion of \rm HIV-1 to CD4+ cells are useful in the treatment of an \rm HIV-1 infected subject. Antibodies against such proteins are also useful for inhibition of \rm HIV-1 infection of \rm CD4+ cells
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N-PSDB; AAW88231.
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03-DEC-1998.

Domain Domain

Key

15-MAR-1999

AAW88231;

RESULT 2 AAW8823

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Length 184;

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HIV-1; CCR5; CCR5-delta32; co-receptor; infection; diagnosis; AIDS;
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            gene therapy; human.
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Best Local Similarity
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                                                                                                                                                                                                                                                                   Beretta A,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADC10142;
                                                               Domain
                                                                                    Domain
                                                                                                         Domain
                                                                                                                              Domain
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 6
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  0;
                                                                                                                                                                                                                                                                                                                                                                                                                                          The present sequence is an inactive human CC (Cys-Cys) chemokine receptor 5 (CCR5), which lacks the last 3 transmembrane regions and the regions involved in G protein-coupling. CCR5 or its cDNA can used to diagnose, treat and/or prevent inflammatory diseases, e.g. rheumatoid arthritis, glomerulonephritis, asthma, idiopathic pulmonary fibrosis and psoriasis, viral infections, especially human immunodefliciency virus type 1 or type 2 (HIV-1 or HIV-2) infection, cancer, atherosclerosis and autoimmune disorders. Subjects that express the inactive receptor have a predisposition, or resistance to HIV-1 and/or HIV-2.
                                                                                                          Inactive; human Cys-Cys chemokine receptor 5; CCR5; human immunodeficiency virus; type 1; type 2; HIV-1; HIV-2; predisposition; resistance; diagnosis; treatment; prevention; inflammatory disease; rheumatoid arthritis; glomerulonephritis; asthma; idiopathic pulmonary fibrosis; psoriasis; viral infection; cancer; atherosclerosis; autoimmune disorder.
                                                                                                                                                                                                                                                                                                                                                                             Active and inactive forms of human CC chemokine receptor CCR-5 - useful to diagnose, prevent and/or treat inflammatory disorders, autoimmune disease and viral infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 .;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         100.0%; Score 164; DB 2; Length 215; 100.0%; Pred. No. 9.5e-17; ive 0; Mismatches 0; Indels (
                                                                                                                                                                                                                                                                                                                             Libert F;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               HIV-1 co-receptor CCR5 variant CCR5-delta32.
                                                                                                                                                                                                                                                                                                                             Vassart G,
                         AAW27408 standard; protein; 215 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAW88238 standard; protein; 215 AA
                                                                                                                                                                                                                                                                                                                                                                                                                      Claim 7; Fig 1d-e; 94pp; English.
                                                                                                                                                                                                                                                                        96EP-00870021.
96EP-00870102.
                                                                                                                                                                                                                                                    97WO-BE000023
                                                                   (first entry)
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                                                                                                                                                                                                                                                                                                                           Parmentier M,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               31; Conservative
                                                                                                                                                                                                                                                                                                       (EURO-) EUROSCREEN SA.
                                                                                       Inactive human CCR5
                                                                                                                                                                                                                                                                                                                                               WPI; 1997-479829/44.
N-PSDB; AAT90118.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 215 AA;
                                                                                                                                                                                     Homo sapiens
                                                                                                                                                                                                       WO9732019-A2
                                                                                                                                                                                                                                                   28-FEB-1997;
                                                                                                                                                                                                                                                                      01-MAR-1996;
06-AUG-1996;
                                                                   14-APR-1998
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                                                                                                                                                                                                                                                                                                                            Samson M,
                                              AAW27408;
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   RESULT 4
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This is the amino acid sequence of a CCR5 variant protein, designated CCR5-delta32, that includes the first 4 transmembrane domains of wild-CCR5-delta32, that includes the first 4 transmembrane domains of wild-CCR5 (see AAW882312), but lacks transmembrane domains 5-7. CCR5 strains of HIV-1. Individuals homozygous for the CCR5-delta32 mutation of the are resistant to HIV-1 infection but heterozygous individuals are susceptible. The invention additionally relates to the identification of variant CCR5m303 (see AAW88231), which lacks transmembrane domains 3-7 of CCR5. The detection of CCR5 variants may be used to identify individuals at lower risk of infection relative to the general population who, if infected, may exhibit slower progression to AIDS. Probes and primers (see AAV84127-36) are provided for use in diagnostic methods for detecting the presence of such variants. A method is provided for inhibiting HIV-1 infection of a cell expressing the CCR5 receptor. This involves introducing a nucleic acid encoding a CCR5 variant into the cell, thereby cells.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                New CCR5 variant protein of the HIV-1 co-receptor - useful in developing resistance of CCR5-expressing cells to HIV-1 infection.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Braun J;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Indels
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100.0%; Pred. No. 9.5e-17;
iive 0; Mismatches 0;
                                                                                                              /note= "transmembrane domain 2"
103. .124
                                                                                                                                                                             103. .124
/note= "transmembrane domain 3"
                                                                       "transmembrane domain 1"
                                                                                                                                                                                                                                                                                                        /note= "transmembrane domain 4"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Quillent C, Arenzana Siesdedos F,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (MOND-) FOND MONDIALE RECH & PREVENTION SIDA.
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Location/Qualifiers
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cytostatic; antidiabetic; anorectic; cerebroprotective; neuroprotective; antiinflammatory; gene therapy; antisense therapy; thyromimetic; NOVX; pathology; cancer; diabetes; obesity; endocrine disorder; CNS disorder; inflammatory disorder; chromosome mapping; tissue typing; predictive medicine.
                                                                                                                                               04-JUN-2001; 2001US-0295607P.

04-JUN-2001; 2001US-0295661P.

06-JUN-2001; 2001US-0296404P.

07-JUN-2001; 2001US-0296418P.

07-JUN-2001; 2001US-0296575P.

11-JUN-2001; 2001US-0295573P.

12-JUN-2001; 2001US-0295573P.

14-JUN-2001; 2001US-029858P.

15-JUN-2001; 2001US-029913P.

19-JUN-2001; 2001US-029913P.

19-JUN-2001; 2001US-029913P.

19-JUN-2001; 2001US-029913P.
                                                                                                                                                                                                                                                                                                                                                                                         2002US-0359122F.
2002US-0359034F.
2002US-0359034F.
2002US-0359121F.
2002US-0359121F.
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2001US-0299133P.
2001US-0299230P.
2001US-029949P.
2001US-0300177P.
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2002US-0379444P.
2002US-00379444.
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2001US-0302951P.
2001US-0308890P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                2002US-0363430P.
2002US-0363676P.
                                                                                                                                                                                                                                                                                          2001US-0300883P.
2001US-0301530P.
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2001US-0324669P.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      WPI; 2003-210149/20.
N-PSDB; ADC10141.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CNS diseases.
                                                                                        WO2003000842-A2
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03-JUL-2001; 3
31-JUL-2001; 3
                                                                                                                                                                                                                                                                                                                                                                                                      22-FEB-2002; 22-FEB-2002; 22-FEB-2002; 2
                                                                                                                                                                                                                                                                                22-JUN-2001;
26-JUN-2001;
28-JUN-2001;
                                                                                                                                                                                                                                                                                                                                                                                                                                    22-FEB-2002;
27-FEB-2002;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       04-JUN-2002;
                                                                                                                              04-JUN-2002;
                                                                     Homo sapiens
                                                                                                           03-JAN-2003
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Agee ML, Anderson DW, Berghs C, Casman SJ, Catterton E;
Dipippo VA, Edinger SR, Eisen A, Ellerman K, Gangolli EA;
Gerlach VL, Gorman L, Guo X, Herrmann JL, Hjalt T, Ji W, Kekuda R;
Khramtson NV, Li L, Liu X, Malyankar UM, Miller CE, Millet I;
Ort T, Padigaru M, Parturajan M, Pena CEA, Rastelli L, Rieger DK;
Rothenberg ME, Shenoy SG, Shimkets RA, Smithson G, Spaderna SK;
Spytek KA, Stone DJ, Vernet CAM, Zhong H, Zhong M, Alsobrook JP; Burgess CE, Lepley DM;

New isolated NOVX polypeptides and nucleic acid molecules useful for treating, preventing and diagnosing pathological conditions with NOVX-associated disorders, such as cancer, obesity, diabetes and inflammatory

Claim 1; SEQ ID NO 162; 772pp; English

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The invention relates to novel isolated polypeptides, mature form of the polypeptide, a sequence that is 95% identical to the polypeptide or the polypeptide comprising one or more conservative substitutions. The NOVX polypeptide is useful for treating or preventing a pathology associated with the polypeptide e.g. disorders associated with abservant expression or activity of the polypeptide, such as cancer, diabetes, obesity, and endocrine. ONS and inflammatory disorders. They can also be used in and predictive medicine. This sequence corresponds to one of the polypeptides of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               cytostatic; antidiabetic; anorectic; cerebroprotective; neuroprotective; antiinflammatory; gene therapy; antisense therapy; thyromimetic; NOVX; pathology; cancer; diabetes; obesity; endoctine disorder; CNS disorder; inflammatory disorder; chromosome mapping; tissue typing; predictive medicine.
                                                                                                                                                                                                                                                         Gaps
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0
                                                                                                                                                                                                                       Query Match
100.0%; Score 164; DB 7; Length 268;
Best Local Similarity 100.0%; Pred. No. 1.2e-16;
Matches 31; Conservative 0; Mismatches 0; Indels (
                                                                                                                                                                                                                                                                                        1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                         1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human NOVX polypeptide SEQ ID NO: 164.
                                                                                                                                                                                                                                                                                                                                                                                                             ADC10144 standard; protein; 268 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         04-JUN-2001; 2001US-0295607P.
04-JUN-2001; 2001US-029561P.
06-JUN-2001; 2001US-029561P.
06-JUN-2001; 2001US-029644P.
07-JUN-2001; 2001US-0296575P.
11-JUN-2001; 2001US-029573P.
12-JUN-2001; 2001US-029573P.
12-JUN-2001; 2001US-02952BE.
18-JUN-2001; 2001US-02982BE.
18-JUN-2001; 2001US-0299139P.
19-JUN-2001; 2001US-0299139P.
22-JUN-2001; 2001US-0299139P.
22-JUN-2001; 2001US-0299139P.
24-JUN-2001; 2001US-0299139P.
25-JUN-2001; 2001US-0301530P.
26-JUN-2001; 2001US-0301530P.
26-JUN-2001; 2001US-0301530P.
27-JUN-2001; 2001US-0301550P.
25-SEP-2001; 2001US-0301550P.
25-SEP-2001; 2001US-0341562P.
21-FEB-2002; 2002US-0358978P.
22-FEB-2002; 2002US-0358978P.
22-FEB-2002; 2002US-0358978P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 04-JUN-2002; 2002WO-US017443
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               18-DEC-2003 (first entry)
                                                                                                                                                                                             Sequence 268 AA;
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us-09-852-238a-5.rag

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The present sequence is human CC (Cys-Cys) chemokine receptor 5 (CCR5), which is stimulated by MIP-1 alpha, MIP-1 beta or RANTES chemokines, but not by monocyte chemoattractant protein 1 (MCP-1), MCP-2, MCP-3, interleukin-8 (IL-8) or growth related gene product alpha (GRO alpha) chemokines. Active CCR-5 is also a receptor of human immunodeficiency virus type 1 or type 2 (HIV-1 or HIV-2). CCR5 or its cDNA can used to diagnose, treat and/or prevent inflammatory diseases, e.g. rheumatoid arthritis, glomerulonephritis, asthma, idiopathic pulmonary fibrosis and psoriasis, viral infections, especially HIV-1 or HIV-2 infection, cancer, atherosclerosis and autoimmune disorders
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Chemokine receptor 88C; atherosclerosis; rheumatoid arthritis; tumour; asthma; viral infection; AIDS; inflammation; autoimmune disease; therapy; diagnosis; leukocyte trafficking; G protein coupled receptor; ligand; modulator; antibody; human.
                                                                                                                                                                                                                                                                                 to diagnose, prevent and/or treat inflammatory disorders, autoimmune disease and viral infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            100.0%; Score 164; DB 2;
100.0%; Pred. No. 1.7e-16;
ative 0; Mismatches 0;
                                                                                                                                                                                                                                                                             Active and inactive forms of human CC chemokine
                                                                                                                                                                                         Vassart G, Libert F;
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125. .145
/label= Intracellular_domain
166. .191
/label= Extracellular_domain
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/label= Extracellular_domain
56. .67
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                                                                                                                                                                                                                                                                                                                                                          Claim 4; Fig 1b-c; 94pp; English
                                                       97WO-BE000023.
                                                                                            96EP-00870021
96EP-00870102
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/label= In
                                                                                                                                                                                       Parmentier M,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  31; Conservative
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                                                                                                                                               (EURO-) EUROSCREEN SA
                                                                                                                                                                                                                        WPI; 1997-479829/44.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Local Similarity
                                                                                                                                                                                                                                            N-PSDB; AAT90117
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 352 AA;
                                                     28-FEB-1997;
                                                                                                              06-AUG-1996;
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                 04-SEP-1997
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     The invention relates to novel isolated polypeptides, mature form of the polypeptide, a sequence that is 95% identical to the polypeptide or the polypeptide comprising one or more conservative substitutions. The Novy polypeptide is useful for treating or preventing a pathology associated with the polypeptide e.g. discorders associated with abstrant expression or activity of the polypeptide, such as cancer, diabetes, obesity, and endocrine. CNS and inflammatory disorders. They can also be used in and predictive medicine. This sequence corresponds to one of the polypeptides of the invention.
                                                                                                                                                                                                                        Agee ML, Anderson DW, Berghs C, Casman SJ, Catterton E;
Dipippo VA, Edinger SR, Eisen A, Ellerman K, Gangolli EA;
Gerlach VL, Gorman L, Guo X, Herrmann JL, Hjalt T, Ji W, Kekuda R;
Khramtsov NV, Li L, Liu X, Malyankar UM, Miller CE, Millet I;
Ort T, Padigaru M, Patturajan M, Pena CEA, Rastelli L, Rieger DK;
Rothenberg ME, Shenoy SG, Shimkets RA, Smithson G, Spaderna SK;
Syrek KA, Stone DJ, Vernet CAM, Zhong H, Zhong M, Alsobrook JP;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       New isolated NOVX polypeptides and nucleic acid molecules useful for treating, preventing and diagnosing pathological conditions with NOVX-associated disorders, such as cancer, obesity, diabetes and inflammatory
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       100.0%; Score 164; DB 7;
100.0%; Pred. No. 1.2e-16;
11ve 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Claim 1; SEQ ID NO 164; 772pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAW27407 standard; protein; 352 AA
                 2002US-0359121P

2002US-035964P

2002US-03608E8P

2002US-0363430P

2002US-0363476P

2002US-0371346P

2002US-0379444
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity 100...
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (first entry)
                                                                                                                                                                                                                                                                                                                                           Stone DJ, r
Lepley DM;
                                                                                                                                                                                         (CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                                  2003-210149/20.
                                                                                                                                                                                                                                                                                                                                                                                                                 N-PSDB; ADC10143.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Sequence 268 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                or CNS diseases,
22-FEB-2002;
22-FEB-2002;
27-FEB-2002;
01-MAR-2002;
                                                                                            12-MAR-2002;
                                                                                                                  10-APR-2002;
                                                                                                                                                 04-JUN-2002;
                                                                            12-MAR-2002;
                                                                                                                                                                                                                                                                                                                                                            Burgess CE,
                                                                                                                                10-MAY-2002
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WO9732019-A2
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAW27407;
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Length 352; Indels

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97WO-US009586.
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                                                                                                                    /label= VI
                                                                                                       .258
                                                                                                                                         .276
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                31; Conservative
                                                                                                                                                                                                                                                                                                                Combadiere C, Feng Y,
Broder CC, Kennedy PE;
                                                                                                                               /note=
                                                                                                                                                                                                                                                                                                                                                  WPI; 1998-032650/03.
N-PSDB; AAT76920.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Sequence 352 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          15-MAR-1999
                                                                                                                                                                                                                                                                        28-MAY-1996;
                                                                                                                                                                                                           W09745543-A2
                                                                                                                                                                                                                              04-DEC-1997.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAW88232;
                                                                                                          Domain
                                                                                                                                          Region
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 Region
                     Domain
                                                                           Domain
                                                     Region
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ID AAW8
XX
XX
AC AAW8
XX
DT 15-M
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ö
                                                                                                                                                                                                                                                                                  This polypeptide sequence comprises novel human chemokine receptor 88C, a give protein coupled receptor that is involved in leukocyte trafficking. Its amino sequence was deduced from a CDNA clone (AAT95161) isolated from a macrophage library. It shows 62% identity to CCCKR1. Chemokine receptor 88-2B (see AAW27124) has also been identified. 88C and 88-2B receptors and their polypeptide fragments can be produced in transformed host cells. The receptors, peptides comprising one or more of the extracellular or intracellular domains, and anti-receptor antibodies can be used to modulate receptor activities, particularly ligand and 6 protein binding, and are potentially potentially useful in the treatment of atheroslerosis, rheumatoid arthritis, tumours, asthma, viral infection, AIDS, inflammatory conditions, pathological immune response,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        CC chemokine receptor 5; CCR5; G-protein coupled receptor;
human immunodeficiency virus; HIV; CD4; AIDS; therapy; transgenic animal.
                                                                                                                                                                                                                      New nucleic acid encoding chemokine receptors 88-2B and 88C - used to modulate leukocyte trafficking, e.g. for treatment of inflammation, tumours, viral infections, auto-immune diseases, etc.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                              ; Score 164; DB 2;
; Pred. No. 1.7e-16;
0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /note= "transmembrane domain"
104. .126
/label= Irin
/note= "transmembrane domain"
               301. .352
/label= Intracellular_domain
   /label= Extracellular_domain
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                        abnormal haematopoietic processes etc
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human CC chemokine receptor 5 (CCR5).
                                                                                                                                                                    Raport CJ;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Location/Qualifiers
                                                                                                                                                                                                                                                                   Claim 16; Page 47-48; 65pp; English
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAW23835 standard; protein; 352 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 100.0%;
100.0%;
                                                                                         96WO-US020759
                                                                                                              95US-00575967
96US-00661393
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/label≖ I
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                                                                                                                                                                    Gray PW, Schweickart VL,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Local Similarity 100.
nes 31; Conservative
                                                                                                                                                                                          NPI; 1997-341689/31.
                                                                                                                                               (ICOS-) ICOS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                             Sequence 352 AA;
                                                                                                                                                                                                      N-PSDB; AAT85161
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Homo sapiens
                                                                                           20-DEC-1996;
                                                                                                                 20-DEC-1995;
                                                 WO9722698-A2
                                                                                                                          07-JUN-1996;
                                                                     26-JUN-1997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAW23835;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
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                   Domain
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Matches
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chemokine receptor that has been designated CCR5. The sequence was deduced from an isolated cDNA clone (see AAT76920). An Ala127Leu variant (see W238340 of CCR5 was also identified. The susceptibility of human macrophages to HIV infection depends on cell surface expression of CD4 and CCR5. CCR5 is a member of the 7-transmembrane superfamily of G-protein coupled cell surface molecules. It plays an essential role in the membrane fusion step of infection by some HIV isolates. The escablishment coexpress human CD4 and CCR5 provides valuable tools for research of HIV infection. Antibodies that bind to CCR5, CCR5 variants, and CCR5-binding agents capable of blocking membrane fusion between HIV and target cells
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             protein sequence comprises of a novel human macrophage-selective CC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          represent potential anti-HIV therapeutics for macrophage tropic strains
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          CC chemokine receptor 5 polypeptide - used to inhibit membrane fusion between HIV and a target cell.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Murphy PM;
                                                                                                                                                                                                                                                                                                                                                                                   /note= "extracellular loop-3 (Claim 19)"
277. .300
'note= "extracellular loop-1 (Claim 19)"
43. 171
'label= IV
                                                                                                                                                     /note= "extracellular loop-2 (Claim 19)"
194. 219
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Berger EA, Alkhatib G,
                                                                                                                                                                                                                 /label= V
/note= "transmembrane domain"
                                                                                                                                                                                                                                                                                                                                       "transmembrane domain"
                                                                                                                                                                                                                                                                                                                                                                                                                                                              /label= VII
/note= "transmembrane domain"
                                                                                 /note= "transmembrane domain"
187. .210
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (USSH ) US DEPT HEALTH & HUMAN SERVICES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAW88232 standard; protein; 352 AA
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New CCR5 variant protein of the HIV-1 co-receptor - useful in developing resistance of CCR5-expressing cells to HIV-1 infection.
                       HIV-1; CCR5; CCR5m303; co-receptor; infection; diagnosis; AIDS;
                                                                    32. .56
/note= "transmembrane domain 1"
                                                                                          /note= "transmembrane domain 2"
                                                                                                                                'note= "transmembrane domain 3"
                                                                                                                                              /note= "transmembrane domain 4"
                                                                                                                                                             "transmembrane domain 5"
                                                                                                                                                                            "transmembrane domain 6"
                                                                                                                                                                                           /note= "transmembrane domain 7"
                                                                                                                                                                                                                                                                                    Arenzana Siesdedos F,
                                                                                                                                                                                                                                                                   (MOND-) FOND MONDIALE RECH & PREVENTION SIDA
                                                                                                        'note= "corresponds to
                                                                                                                                                                                                                                                                                                                                            Disclosure; Page 34-35; 55pp; English.
                                                             Location/Qualifiers
                                                                                                                 in CCr5m303"
                                                                                                                                                                                                                                      98WO-EP003437
                                                                                                                                                                                                                                                     97US-0048057P
                                                                                                                                       .167
                                                                                                                                                                    .260
                                                                                                                         .124
                                                                                                                                                     . 223
                                                                                                                                                                                    .301
                                                                                    .87
        HIV-1 co-receptor CCR5.
                                                                                                                                                                                                                                                                                  Quillent C,
                                                                                                                                                             /note=
                                                                                                                                                                            note=
                                                                                                                 (Stop)
                              gene therapy; human.
                                                                                                                                                                                                                                                                                                 1999-059835/05.
                                                                                                                                                                                                                                                                                                       N-PSDB; AAV84126.
                                                                                                Misc-difference
                                              Homo sapiens
                                                                                                                                                                                                       WO9854317-A1
                                                                                                                                                                                                                                       29-MAY-1998;
                                                                                                                                                                                                                                                     30-MAY-1997;
                                                                                                                                                                                                                                                                                   Beretta A,
                                                             Key
Domain
                                                                                   Domain
                                                                                                                       Domain
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This is the amino acid sequence of wild-type human CCRS, which serves as a co-receptor for infection by macrophage-tropic (M-tropic) strains of HIV-1. The invention relates to the identification of a CCRS variant (see AAM88231), designated CCRSA303, comprising the first two transmembrane domains of wild-type CCRS, but lacking transmembrane domains 3-7. The presence of the CCRSA303 variant with the wild type CCRS allele shows a positive correlation with resistance to infection with M-tropic HIV-1 strains, and may indicate slower progression of the disease. The cits of infection relative to the general population who, if infected may exhibit slower progression to ALDS. Probes and primers (see AAV884127-36) are provided for use in diagnostic methods for detecting the presence of such variants. A method is provided for inhibiting HIV-1 infection of a cell expressing the CCRS receptor. This involves introducing a mucheic acid encoding a CCRS variant into the cell, thereby reducing the number of functional CCRS molecules present on the cell surface

Sequence 352 AA;

Gaps 0; Length 352; 0; Indels 100.0%; Score 164; DB 2; 100.0%; Pred. No. 1 7e-16; Mismatches · 0 Ouery Match
Best Local Similarity 100.

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1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31

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human immunodeficiency virus, antimicrobial; vasodilator; vulnerary, cytostatic; immunosuppressive; noctropic; neuroprotective; gene therapy; neurodegenerative disorder; Kaposi's sarroma; autoimmune disease; rheumatoid arthritis; cancer; breast; ovary; adrenal gland; bone marrow; gastrointestinal tract; lung; liver; immune disorder; Addison's disease; haemolytic anaemia; autoimmune thyroiditis; diabetes mellitus; allergy; multiple sclerosis; ulcerative colitis; Crohn's disease; wound healing; cardiovascular disorder; myocardial ischaemia. Human; G-protein chemokine receptor; CCR5; HDGNR10; inflammation; HIV; Human G-protein chemokine receptor (CCR5) HDGNR10 protein #2. AAE07048 standard; protein; 352 AA (first entry) WO200158916-A2 Homo sapiens. 16-OCT-2001 16-AUG-2001. AAE07048; RESULT 12 AAE07048 

TGT (Cys) in wild-type CCR5, TGA

09-FEB-2001; 2001WO-US004153. 09-FEB-2000; 2000US-0181258P. 09-MAR-2000; 2000US-0187999P.

09-MAR-2000; 2000US-0187999P. 22-SEP-2000; 2000US-0234336P.

(HUMA-) HUMAN GENOME SCI INC

Li Y, Ruben SM; Rosen CA, Roschke V,

Braun J;

WPI; 2001-488966/53. N-PSDB; AAD13299 Isolated nucleic acid encoding a human G-protein chemokine receptor (CCR5) HDGNR10 polypeptide, useful for preventing or treating autoimmune diseases e.g. rheumatoid arthritis, hyperproliferative disorders and neurodegenerative disorders.

Example 40; Page 504-505; 518pp; English

The invention relates to human G-protein chemokine receptor (CCRS)

Useful for treating, preventing or ameliorating a disease or disorder

useful for treating, preventing or ameliorating a disease or disorder

associated with inflammation, defective or aberrant chemotaxis of immune

cells, HIV infection (such as Phemocystis carinii pneumonia or Kaposi's

sarcoma) or defective or aberrant T-cell antigen presenting cell

interaction. The disease or disorder may also be an infectious disease

(e.g. a vixal infection, or a poxvirus infection), an autoimmune

clisease (e.g. rheumatoid arthritis) or a neurodegenerative disorder. The

disease or disorder may be associated with aberrant CCRS expression, lack

cutofine, cCRS function, aberrant CCRS ligand expression, or lack of CCRS ligand

cutofine. CCRS function protein is used as a food additive or preservative

con increase or decrease storage capabilities. CCRS HDGNRIO DNA, are useful

con chromosome identification and in gene therapy. CCRS HDGNRIO DNA,

controlly antibodies, agonists and antagonists are also useful in the

dispanse or dispanse treatment and prevention of cancer (breast, ovary, adrenal) haemolytic anaemia, autoimmune thyroiditis, diabetes mellitus, Crohn's disease, multiple sclerosis, rheumatoid arthritis and ulcerative colitis); cardiovascular disorders (myocardial ischaemias) and wound healing. The present sequence is human CCRS HDGNR10 protein gland, bone, bone marrow, gastrointestinal tract, liver, lung, urogenital); immune disorders (Addison's disease, allergies, autoimmune haemolytic anaemia,

Sequence 352 AA;

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This invention describes a novel diagnostic agent (A) comprising at least two different ligands (I) for receptors (II) that are implicated in disease. (A) are used for the diagnosis of tumors (especially colorectal or prostatic), organ rejection, inflammation and autoimmume diseases.

Cor prostatic), organ rejection, inflammation and autoimmume diseases.

Also inhibitors of (I) are used therapeutically against tumors (and their metastases), inflammation (particularly bronchial atthma or chronic bowel inflammation), or autoimmume diseases (Theumatoid arthritis or lupus), where the (cardio) vascular, lymphatic, respiratory, nervous, diseases.

Comparison diseases The products of the invention are enfected, and bone marrow diseases. The products of the invention are chemokine derivatives which have cytostatic, antiinflammatory, antiathmatic, immunosuppressive, dermatological, antirheumatic, antiarthritic.

Commonshines act on specific tumor and inflammatory calls through a constellation of chemokine receptors (CR), which control migration and constellation of these cells. Ads00128 represent human chemokine fragments used to illustrate the method of the invention
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Diagnostic agent containing two or more receptor-specific ligands, useful for detecting tumors, inflammation etc., also therapeutic use of ligand inhibitors.
                                                                                                                                                                                                                                                                                                                                                      Chemokine; tumour diagnosis; colorectal; prostatic; organ rejection; inflammation; autoimmune disease; metastasis; bronchial asthma; lupus; chronic bowel inflammation; rheumatoid arthritis; cytostatic; antiinflammatoxy; antiasthmatic; immunosuppressive; dermatological; antirheumatic; antiarthritic.
                                               Gaps
                                               0
           Length 352;
                                               0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Spodsberg
             Score 164; DB 4;
Pred. No. 1.7e-16;
100.0%; Scor.
100.0%; Pred. No. 1.
0; Mismatches
                                                                                     31
                                                                                                                     1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                     1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Disclosure; Page 10; 26pp; German.
                                                                                                                                                                                                                    AAG80111 standard; protein; 352 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    31-MAR-2000; 2000DE-01016013.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      02-APR-2001; 2001WO-EP003708.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Forssmann W, Adermann K,
                                                                                                                                                                                                                                                                                          (first entry)
 Query Match
Best Local Similarity luv.
Local 31; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PHARM GMBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 WPI; 2001-626256/72
                                                                                                                                                                                                                                                                                                                                Human CCR5 protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (FORS/) FORSSMANN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 352 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Homo sapiens.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (IPFP-) IPF
                                                                                                                                                                                                                                                                                            17-JAN-2002
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                                                                                                                                                                                RESULT 13
AAG80111
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The present invention relates to a transformed mammalian cell that contains a gene encoding CD4, a construct encoding a reporter gene under the regulation of an human immuno deficiency virus (HIV) long terminal crepeat (ITR) and that has been transduced with a vector encoding a human chemokine receptor (CKR) where the CD4 and the CKR are present on the coll surface of transformed mammalian cell. The invention is useful for identifying drugs or antibodies that interfere with the translocation of HIV into transformed mammalian cell or for identifying a human chemokine creeptor that facilitates the infection of a particular HIV strain into the transformed mammalian cell. Compounds identified can be used to treat cellular dysfunction and to prevent or combat HIV infection. The present sequence is a human chemokine receptor (CKR), CC-CKR-5 related protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 mammalian cell (I) that contains a CD4 gene, reporter gene for identification of drugs and antibodies for treatment of
                                                                                                                                                                                                         Human, transformed mammalian cell; CD4; reporter gene; translocation; human immuno deficiency virus; HIV; long terminal repeat; LTR; therapy; chemokine receptor; CKR; cellular dysfunction; HIV infection; cofactor;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ;
0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 352;
                                                                                                                                                                                Human chemokine receptor (CKR), CC-CKR-5 related protein #2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Liu R;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          100.0%; Score 164; DB 4; 100.0%; Pred. No. 1.7e-16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Landau NR,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          0; Mismatches
envelope glycoprotein; anti-HIV.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Disclosure, Col 47-50; 37pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (AARO-) AARON DIAMOND AIDS RES CENT.
(UYNY ) UNIV NEW YORK STATE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Ellmeier W,
                                                                                            Ā
                                                                                            AAE04321 standard; protein; 352
                                                                                                                                                                                                                                                                                                                                                                                      97US-00861105.
                                                                                                                                                                                                                                                                                                                                                                                                                     96US-0017157P.
                                                                                                                                                                                                                                                                                                                                                                                                                                     96US-0020043P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                   97US-00858660.
                                                                                                                                                         (first entry)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             31; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Littman DR, Deng H,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             2001-417127/44.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 352 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WPI; 2001-417127,
N-PSDB; AAD08577
                                                                                                                                                                                                                                                                                                 Homo sapiens.
                                                                                                                                                                                                                                                                                                                              US6258527-B1.
                                                                                                                                                                                                                                                                                                                                                                                        21-MAY-1997;
                                                                                                                                                                                                                                                                                                                                                                                                                                      19-JUN-1996;
19-MAY-1997;
                                                                                                                                                                                                                                                                                                                                                                                                                        20-MAY-1996;
                                                                                                                                                         04-SEP-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Transformed
and HIV LTR
                                                                                                                                                                                                                                                                                                                                                            10-JUL-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                                                    CC-CKR-5;
                                                                                                                            AAE04321;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Local
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Matches
                                                                   RESULT 14
                                                                                   AAE04321
                                                                                                              a
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similarity 100.0%; Score 164; DB 4; Length 352; Similarity 100.0%; Pred. No. 1.7e-16; Conservative 0; Mismatches 0; Indels

RESULT 15 AAE07039

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Gaps

.; 0

.. 0

31; Conservative

Query Match Best Local 9 Matches თ

16-OCT-2001 AAE07039 

AAE07039 standard; protein; 352 AA.

(first entry)

Human G-protein chemokine receptor (CCR5) HDGNR10 protein #2.

human immunodeficiency virus, antimicrobial, vasodilator; vulnerary, cytostatic; immunosuppressive; antimicrobial; vasodilator; vulnerary, neurodegenerarive disorder; Kaposi s arcoma; autofmmune disease; rheunatoid arthritis; cancer; breast; ovary, adrenal gland; bone marrow; gastrointestinal tract; lung; liver; immune disorder; Addison's disease; multiple sclerosis; ulcerative collitis; diabetes mellitus; allergy; authibe sclerosis; ulcerative collitis; Crohn's disease; wound healing; cardiovascular disorder; myocardial ischaemia. Human; G-protein chemokine receptor; CCR5; HDGNR10; inflammation; HIV;

Homo sapiens.

WO200158915-A2.

16-AUG-2001

09-FEB-2001; 2001WO-US004152.

09-FEB-2000; 2000US-0181258P. 09-MAR-2000; 2000US-0187999P. 22-SEP-2000; 2000US-0234336P.

(HUMA-) HUMAN GENOME SCI INC

Ruben SM; Li Y, Roschke V, Rosen CA,

WPI; 2001-488965/53 N-PSDB; AAD13198 Isolated nucleic acid encoding a human G-protein chemokine receptor (CCR5) HDGNR10 polypeptide, useful for preventing or treating autoimmune diseases e.g. rheumatoid arthritis, hyperproliferative disorders and neurodegenerative disorders.

Example 40; Page 486-487; 495pp; English.

protein. CORS HOGNRIO antibodies are useful for treating, preventing or ameliorating a disease or disorder associated with inflammation, defective or aberrant chemotaxis of immune cells, HIV infection (such as pneumocystis carinii pneumonia or Kaposi's sarcoma) or defective or disorder may also be an infections degrace. The disease or an early stage HIV infection, a cytomegalovirus infection such as an early stage HIV infection, a cytomegalovirus infection or a poxvirus infection), an autoimmune disease (e.g. rheumatoid arthritis) or a neurodegenerative disorder. The disease (e.g. rheumatoid arthritis) or a neurodegenerative disorder. The disease or disorder may be associated with aberrant CRS expression, lack of CRS function. GCRS HDGNRIO protein is used as a food additive or preservative to increase or decrease storage capabilities. CRS HDGNRIO DNA are useful for chromosome identification and in gene therapy. CRS HDGNRIO DNA, protein, and antagonists are also useful in the diagnosis, treatment and an arthribus and antagonists are also useful in the diagnosis, The present sequence is human G-protein chemokine receptor (CCRS) HDGNR10 treatment and prevention of cancer (breast, ovary, adrenal gland, bone, bone marrow, gastrointestinal tract, liver, lung, urogenital); immune disorders (Addison's disease, allergies, autoimmune haemolytic anaemia, autoimmune thyroiditis, diabetes mellitus, Crohn's disease, multiple sclerosis, rheumatoid arthritis and ulcerative colitis); cardiovascular disorders (myocardial ischaemias) and wound healing

Sequence 352 AA;

0; Gaps ; 100.0%; Score 164; DB 4; Length 352; 100.0%; Pred. No. 1.7e-16; ive 0; Mismatches 0; Indels 0 Query Match
Best Local Similarity 100.
Matches 31; Conservative

31 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR g à

completed: July 29, 2004, 13:28:33 53 secs Search co

(0102N) AMDIA 8604 2141

GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd.

- protein search, using sw model OM protein July 29, 2004, 13:26:01; Search time 16 Seconds (without alignments) 186.371 Million cell updates/sec Run on:

US-09-852-238A-5

164 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31 Title: Perfect score: Sednence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

283366 segs, 96191526 residues Searched:

Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 su

PIR 78:\*
1: pir1:\*
2: pir2:\*
3: pir3:\*
4: pir4:\* Database :

summaries

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

50	Description	chemokine (C-C) re	hypothetical prote	actin depolymerizi	protein-tyrosine k	uracil phosphoribo	5-methyltetrahydro	5-methyltetrahydro	actin depolymerizi	conserved hypothet	E2 glycoprotein pr	hypothetical prote	probable exported	E2 glycoprotein -	E2 glycoprotein pr	E2 glycoprotein -	spike protein chai	E2 glycoprotein pr			E2 glycoprotein pr	E2 glycoprotein pr	laminin beta-1 cha	hypothetical prote	protein F59B10.1 [	chemokine (C-C) re	chemokine (C-C) re	actin-depolymerizi	٤	hypothetical prote
SUMMARIES	ID	A43113	T20317	T47540	PT0189	B86864	AF1284	AI1655	G84717	T40181	VGIHAK	T40491	AE0169	S14599	S14600	S14598	S41626	VGIHD6	T29089	VGIHIB	S14939	S14940	MMFFB1	T22982	H88293	JC2443	I38450	S30934	S	136810
	DB	7	~1	7	7	~	7	7	7	N	Н	N	0	7	7	Ø	7	М	~	-	~	7	н	N	71	~	7	7	ď	7
	Length	352	178	130	54	211				516	1162	818	364	520	520	520	544	550	1130	1154	1162	1162	1790	1039	1192	360	374	126	130	141
ok	Query Match	100.0	34.8	33.2	32.9	32.9	32.9	32.9	32.6	32.3	31.7	31.4	31.1	31.1	31.1	31.1	31.1	31.1	31.1	31.1	31,1	31.1	31.1	30.2	30.2	29.9	29.9	29.6	29.6	29.6
	Score	164	57	54.5	54	54	54	S	53.5	53	52	51.5	51	51	51	51	51	51	51	51	51	51	51	49.5	49.5	49	4.9	48.5	48.5	48.5
	Result No.		5	ю	4	Ŋ	9	7	80	6	10	11	12	13	14	15	16	17	18	1-9	20	21	22	23	2.4	25	26	27	28	29

chemokine (C-C) re	decorin binding pr	hypothetical prote	probable membrane	actin-depolymerizi	actin-depolymerizi	conserved hypothet	outer membrane pro	probable outer mem	hypothetical prote	G protein-coupled	hypothetical prote	hypothetical prote	galactosamine-cont	protein-tyrosine k	hypothetical prote
A45177	H70209	T23559	861116	T02883	T02882	H82265	F64599	C71912	S48954	S55594	T33464	S50452	E69631	S23251	F90099
7	7	~	7	~	7	N	~	~	7	~	7	~	7	7	7
355	191	581	612	139	139	258	305	307	378	383	528	999	900	942	1005
29.6	29.3	29.3	29.3	29.0	29.0	28.7	28.7	28.7	28.7	28.7	28.7	28.7	28.7	28.7	28.7
48.5	48	48	48	47.5	47.5	47	47	47	47	47	47	47	47	47	47
30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

RESULT 1

283366

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				0-Jun		Σ.		ian CC.		
				96 #text_change 20	C; Accession: A43113; S71808; A58834; A58832; G02653; A58833	R; Samson, M.; Labbe, O.; Mollereau, C.; Vassart, G.; Parmentier, M.	*** *****	A;TITLE: Molecular cloning and functional expression of a new human CC-ch	8639485	
	lan	RS		vision 12-Jul-19	1834; A58832; G02	au, C.; Vassart,	. 4	unctional expres	):96241590; PMID:	The second secon
	sceptor 5 - hum	: C-C CKR-5; CC	npiens (man)	96 ∦sednence re	[3; S71808; A58	De, O.; Mollere	3362-3367, 1996	cloning and f	r: A43113; MUID	. 61
A43113	chemokine (C-C) receptor 5 - human	N;Alternate names: C-C CKR-5; CCR5	C;Species: Homo sapiens (man)	;Date: 12-Jul-199	Accession: A431	Samson, M.; Lab	lochemistry 35,	Title: Molecula	A; Reference number: A43113; MUID: 96241590; PMID: 8639485	Accession: A431
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A; Molecule type: mRNA A; Residues: 1-352 <SAM1>

A,Cross-references: GB:X91492; NID:g1262810; PIDN:CAA62796.1; PID:g1262811
R,Samson, M.; Libert, F.; Doranz, B.J.; Rucker, J.; Liesnard, C.; Farber, C.M.; Saragosti M.; Imai, T.; Rana, S.; Yi, Y.; Smyth, R.J.; Collman, R.G.; Doms, R.W.; Vassart, G.; Pan Nature 382, 722-725, 1596
A,Title: Resistance to HIV-1 infection in caucasian individuals bearing mutant alleles of A;Reference number: S71808; MUID:96345670; PMID:8751444

A;Status: nucleic acid sequence not shown; not compared with conceptual translation A;Molecule type: DNA A;Residues: 182-206;207-230 <SAM2> A;Residues: 182-206;207-230 <SAM2> A; Accession: S71808

A,Status: nucleic acid sequence not shown; not compared with conceptual translation A,Status: nucleic acid sequence not shown; not compared with conceptual translation A,Status: nucleic acid sequence not shown; not compared with conceptual translation A,Molecule type: DNA A,Residues: 1-184, IKDSHIGAGPAAACHGHILIGNPRNSASVSK <SAM3>
A,Cross-references: GB.X99393; NID:g1524062; PIDN:CAAA7767.1; PID:g1524063
A,Note: this frameshift mutation results in ann-functional receptor but confers a degree of may have had a selective advantage by conferring resistance to Yersinia plague infects R; Combadiere, C.; Ahuja, S.K.; Tilffany, H.L.; Murphy, P.M.

W. Leukoc, Biol. 60, 147-152, 1996
A,FtTe: Cloning and functional expression of CC CRR5, a human monocyte CC chemokine recent annear name and functional expression of CC CRR5, a human monocyte CC chemokine recent annear name and functional expression. As 8993
A,FtTe: Cloning and functional expression of CC CRR5, a human monocyte CC chemokine recent annear name and annear name annear name

submitted to the EMBL Data Library, May 1996 A, Reference number: H01541

A;Accession: G02653
A;Status: translated from GB/EWBL/DDBJ
A;Molecule type: mRNA
A;Molecule type: mRNA
A;Residues: 1-89, Lv. 91-352 <COM2>
A;Cross-references: EMBL:U57840
A;Cross-references: EMBL:U57840
A;Cross-references: EMBL:U57840
A;Cross-references: EMBL:U57840
A;Cross-references: EMBL:U57840
A;Cross-references: A;Cross-reference and functional characterization of a novel human CC chemokine A;Reference number: A58833; MUID:96291862; PMID:8663314

a

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C;Superfauily: unassigned Ser/Thr or Tyr-specific protein kinases; protein kinase homolog
C;Keywords: ATP; phosphotransferase; tyrosine-specific protein kinase
F;1-54/Domain: protein kinase homology (fragment) <KIN>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  uracil phosphoribosyltransferase (EC 2.4.2.9) [imported] - Lactococcus lactis subsp. lact
C;Species: Lactococcus lactis subsp. lactis
C;Date: 23-Mar_2001 #sequence_revision 23-Mar-2001 #text_change 03-Aug-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          R;Bolotin, A.; Wincker, P.; Mauger, S.; Jaillon, O.; Malarme, K.; Weissenbach, J.; Ehrlic
Benome Res. 11, 731-753, 2001
A;Title: The complete genome sequence of the lactic acid bacterium Lactococcus lactis ssg
A;Reference number: A86625; MUID:21235186; PMID:11337471
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                A;Residues: 1-211 <STO>
A;Cross-references: GB:AE005176; PID:g12724952; PIDN:AAX06012.1; GSPDB:GN00146
A;Experimental source: strain IL1403
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         protein-tyrosine kinase (BC 2.7.1.112) tyro-7 - rat (fragment)
C;Species: Rattus norvegicus (Norway rat)
C;Species: Dec-1991 #sequence_revision 31-Dec-1991 #text_change 04-Feb-2000
C;Accession: Pr0189
R;Lai, C.; Lemke, G.
Neuron 6, 691-704, 1391
A;Title: An extended family of protein-tyrosine kinase genes differentially A;Reference number: Pr0183; MUID:91222560; PMID:2025425
A;Accession: P70189
                                                         A;Reference number: 224468
A;Accession: T47540
A;Accession: T47540
A;Accession: T47540
A;Accession: T47540
A;Residues: 1-130 «JOR»
A;Residues: 1-130 «JOR»
A;Cross-references: EMBL;AL162459
A;Experimental source: cultivar Columbia; BAC clone F16L2 C;Genetics:
                                   submitted to the Protein Sequence Database, March 2000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        A;Gene: upp
C;Superfamily: uracil phosphoribosyltransferase upp
C;Keywords: glycosyltransferase; pentosyltransferase
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   2 DYQVSSP-----IYDINYYTSEPCQKINVKQIA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 54.5; Di
Pred. No. 1.5;
5; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |::| ||::|| || DFGLSKKIYNGDYYRQGPFAKMPVKWIA 41
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            2 DYQVSSPIYDINYYTSEPCQKINVKQIA 29
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           32.9%; Score 54; 42.9%; Pred. No. (
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           A; Experimental source: sciatic nerve
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  33.2%;
37.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          13; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Conservative
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Best Local Similarity
Matches 13; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
Matches 12; Conserv
                                                                                                                                                                                                                                                                                                                                                                             A;Introns: 81/2
A;Note: F16L2.210
C;Superfamily: cofilin
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A; Molecule type: mRNA
A; Residues: 1-54 <LAI>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               A, Status: preliminary
A, Molecule type: DNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          C;Accession: B86864
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                                                                                                                                                                                                                                                                                                                                             A; Map position: 3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Gene: tyro-7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           46
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
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A; Molecule type: mRNA
A; Residues: 1-312 <RNA
A; Conment: This is a receptor for chemokines MIP-lalpha (see PIR.30574), MIP-lbeta (see C; Comment: Macrophage- and dual-tropic strains of HIV-1 bind to a complex of chemokine (C; Gametics: GB: US-30510; OMIM: 601373
A; Gane: GDB: CMKBR5; CCR5; CCR-5; CCCCR-5; CCR5; ChemR13
A; Gane: GDB: CMKBR5; CCR5; CCR-5; CCCCR7-5; CRR5; ChemR13
A; Map position: 3p21-3p21
A; Map position: 3p21-3p21
A; Map position: G protein-coupled receptor for chemokines MIP-lalpha, MIP-lbeta and RANTH A; Deterniption: G protein-coupled receptor; glycoprotein; phosphoprotein; transmembrane #status predicted <TM2>
C; Superfamily: vertebrate rhodopsin
C; Superfamily: vertebrate stocoupled receptor; glycoprotein; phosphoprotein; transmembrane #status predicted <TM2>
F; 103-124/Domain: transmembrane #status predicted <TM3>
F; 142-166/Domain: transmembrane #status predicted <TM4>
F; 193-218/Domain: transmembrane #status predicted <TM4>
F; 225-257/Domain: transmembrane #status predicted <TM5>
F; 225-257/Domain: transmembrane #status predicted <TM5>
F; 225-257/Domain: transmembrane #status predicted <TM5>
F; 225-257/Domain: transmembrane #status predicted <TM6>
F; 225-257/Domain
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  actin depolymerizing factor 2 - Arabidopsis thaliana
N;Alternate names: protein F16L2.210
S;Species Arabidopsis thaliana (mouse-ear cress)
C;Date: 20-Apr-2000 #sequence_revision 20-Apr-2000 #text_change 26-May-2000
C;Accession: T47540
R;Jordan, N.; Bangert, S.; Wiedelmann, R.; Voss, H.; Unseld, M.; Mewes, H.W.; Rudd, S.;
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A;Experimental source: clone D1081
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                hypothetical protein D1081.6 - Caenorhabditis elegans
C;Species: Caenorhabditis elegans
C;Species: Caenorhabditis elegans
C;Accession: T20317
R;Dobson, R.
submitted to the EMBL Data Library, July 1996
A;Reference number: Z19256
A;Reference number: Z19256
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      F;20-269,101-178/Disulfide bonds: #status predicted
F;268/Binding site: carbohydrate (Asn) (covalent) #status predicted
F;336,337,342/Binding site: phosphate (Ser) (covalent) #status predi
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0
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Best Local Similarity
'Local 9; Conserve
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Matches 31; Conserv
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A;Introns: 95/3; 114/3
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A,Gene: CESP:D1081.6
A,Map position: 1
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genes differentially expressed in

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Indels

Length 130;

2; 10; 29

DB

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Gaps

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11; Indels

Length 54;

DB 2;

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E2 glycoprotein precursor - avian infectious bronchitis virus (strain KB8523)
RyAlternate names: peplomer protein; spike protein
NyAlternate names: peplomer protein; spike protein
NyContains: E2 glycoprotein subunit S1; E2 glycoprotein subunit S2
C;Species: avian infectious bronchitis virus, IBV
C;Date: 30-Sep-1990 #sequence_revision 30-Sep-1990 #text_change 16-Jul-1999
C;Accession: B29249
R;Sutou, S.; Sato, S.; Okabe, T.; Nakai, M.; Sasaki, N.
Virology 165, S89-595, 1988
A;Title: Cloning and sequencing of genes encoding structural proteins of avian infectious A;Reference number: A29249; MUID:88306251; PMID:2841803
A;Reference number: R29249
A;Molecule type: genomic RNA
                actin' depolymerizing factor 6 [imported] - Arabidopsis thaliana actin' depolymerizing factor 6 [imported] - Arabidopsis thaliana (mouse-ear cress)
C;Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 02-Feb-2001 #sequence_revision 02-Feb-2001 #text_change 16-Feb-2001
C;Accession: G84717
R;Inn, X; Kaul, S; Rounsley, S.D.; Shea, T.P.; Benito, M.I.; Town, C.D.; Fujii, C.Y.; h
N; Koo, H.; Moffat, K.S.; Cronin, L.A.; Shen, M.; VanAken, S.E.; Umayam, L.; Tallon, L.;
Musture, A.D.; Nierman, W.C.; White, O.; Eisen, J.A.; Salzberg, S.L.; Fraser, C.M.; Venter, J.
Nature 402, 761-768, 1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        conserved hypothetical protein SPBC30D10.15 - fission yeast (Schizosaccharomyces pombe)
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C;Date: 03-Dec-1999 #sequence_revision 03-Dec-1999 #text_change 03-Dec-1999
C;Accession: T40181
R;Wood, V; Rajandream, M.A.; Barrell, B.G.; Lauber, J.; Hilbert, H.; Duesterhoeft,
gubmitted to the EMBL Data Library, February 1998
A;Reference number: 221910
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Nolecule type: DNA
A;Residues: 1-516 <WOO>
A;Residues: EMBL:297992; PIDN:CAB10810.1; GSPDB:GN00067; SPDB:SPBC30D10.15
A;Experimental source: strain 972h-; cosmid c30D10
                                                                                                                                                                                                                                       A,Title: Sequence and analysis of chromosome 2 of the plant Arabidopsis thaliana.
A,Reference number: A84420; MUID:20083487; PMID:10617197
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Pred. No. 2.2;
5; Mismatches
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A;Gene: SPDB:SPBC30D10.15
A;Map position: 2
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nes 10; Conserv
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Best Local Similarity
Matches 11; Conserv
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A,Map position: 2
C,Superfamily: cofilin
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                                                                                                                                                                                                                                                                                                                                                                          A; Molecule type: DNA
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C;Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 27-Nov-2001
C;Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 27-Nov-2001
C;Datession: Al1655
C;Gatession: Al1655
C;Gatession: Al1655
D; Jones, L.M.; Karst, U.
D; Jones, L.M.; Karst, U.
D; Jones, L.M.; Karst, U.
Science 294, 849-852, 2001
A;Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Mayathors: Kreft, J.; Kuhn, M.; Tillerrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland, A;Reference number: AB1077; MUID:21537279; PMID:11679669
A;Accession: Al1655
                                                                                                                                                                                                                                                                                                                               C;Species: Listeria monocytogenes
C;Species: Listeria monocytogenes
C;Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 27-Nov-2001
C;Accession: AF1284
C;Accession: AF1284
R;Glaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloecker, D.; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H.; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H. Science 294, 849-882, 2001
A;Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Makathors: Kreft, J.; Kimoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland, A;Fitle: Comparative genomics of Listeria species.
A;Fitle: Comparative genomics of Listeria species.
A;Feference number: AB1077; MUID:21537279; PMID:11679669
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        5-methyltetrahydrofolate-homocysteine methyltransferase (metH) homolog lin1786 [imported
                                                                                                                                                                                                                                                                                          5-methyltetrahydrofolate-homocysteine methyltransferase (metH) homolog lmo1678 [imported
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ij
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10;
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     DB 2;
                                                                                                           1 MDYQVSS--PIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                 MAYEVSRDLPLEDVEIET--PVOKTTVKOIAGK 70
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  Score 54; DB 2
Pred. No. 3.1;
4; Mismatches
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Pred. No.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Experimental source: strain EGD-e
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 32.9%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match 32.9%;
Best Local Similarity 42.3%;
Matches 11; Conservative
  32.9%;
milarity 45.5%;
Conservative 4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Conservative
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Best Local Similarity
Matches 11; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      A; Molecule type: DNA
A; Residues: 1-617 < GLA>
                                    Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Status: preliminary
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        A; Accession: AF1284
A; Status: preliminary
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       C,Genetics:
A,Geme: lin1786
                                                                 15;
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           Query Match
Best Local S:
Matches 15
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us-09-852-238a-5.rpr

Query Match
Best Local Similarity 37.5%;
Matches 9; Conservative 4

⋩ g T40491

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RESULT 13
S14599
E. glycoprotein - avian infectious bronchitis virus (strain UK/142/86) (fragment)
E. glycoprotein - avian infectious bronchitis virus, IBV
N.Alternate names: spike glycoprotein chain S1
C.Species: avian infectious bronchitis virus, IBV
A.Variety: strain UK/142/86
C.Actes 19-Mar-1997 #sequence_revision 10-Oct-1997 #text_change 20-Sep-1999
E.Cavanagh, D.; Davis, P.J.; Cook, J.K.A.; Li, D.; Kant, A.; Koch, G.
E.Cavanagh, D.; Davis, P.J.; Cook, J.K.A.; Li, D.; Kant, A.; Koch, G.
E.Cavanagh, D.; Davis, Data Library, March 1991
A.Description: Infectious bronchitis virus: the S1 spike glycoprotein subunits of some strains.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                glycoprotein precursor - avian infectious bronchitis virus (strain UK/123/82) (fragme
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           site: carbohy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      R;Cavanagh, D.; Davis, P.J.; Cook, J.K.A.; Li, D.; Kant, A.; Koch, G. submitted to the EMBL Data Library, March 1991
A;Description: Infectious bronchitis virus: the S1 spike glycoprotein subunits of some A;Reference number: S14598
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ·,
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N/Alternate names: spike glycoprotein chain S1
C;Species: avian infectious bronchitis virus, IBV
A;Variety: strain UK/167/84
C;Date: 19-Mar-1997 #sequence_revision 19-Mar-1997 #text_change 20-Sep-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               A)Cross-references: EMBL:XB8067; NID:g59029; PIDN:CAA41098.1; PID:g59030 C;Superfamily: coronavirus E2 glycoprotein
C;Keywords: glycoprotein; peplomer protein; spike protein
F;1-520/Product: E2 glycoprotein subunit S1 #status predicted <GS1>F;5,84,121,127,146,161,195,220,247,254,259,262,289,408,430,496,513/Binding
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            A Status: preliminary
Abdlecule type: genomic RNA
A;Residues: 1-520 <CAV>
A;Cross-references: BMBL:X58066; NID:959027; PIDN:CAA41097.1; PID:959028
C;Superfamily: coronavirus E2 glycoprotein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
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46.7%; Pred. No. 22;
tive 3; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              N.Alternate names: spike glycoprotein chain S1
C.Species: avian infectious bronchitis virus, IBV
A.Variety: strain UK/123/82
C.Date: 19-Mar-1997 #sequence_revision 21-Nov-1998
C.Accession: S14600
R.Cavanagh, D.; Davis, P.J.; Cook, J.K.A.; Li, D.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 51; DB 2
Pred. No. 22;
3; Mismatches
                                         239 VDSKVSSIVYDVNSYTDK 256
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           464 YGLNYYKVNPCEDVN 478
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            A; Reference number: S14598
A; Accession: S14599
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Local Similarity
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Matches
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S14600
E2 qlyc
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                                                               C, Superfamily: coronavirus E2, glycoprotein, c) Keywords: glycoprotein, peplomer protein, spike protein F11-18/Domain: signal sequence #status predicted <SIG> F19-537/Product: E2 glycoprotein subunit S1 #status predicted <GS1> F;538-1162/Product: E2 glycoprotein subunit S2 #status predicted <GS2> F;538-1162/Product: E2 glycoprotein subunit S2 #status predicted <GS2> F;51,77,103,144,163,178,212,237,247,264,271,276,283,306,425,447,513,530,579,591,669,676,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  probable exported protein YPO1387 [imported] - Yersinia pestis (strain C092)
C.Species: Versinia pestis
C.Species: Versinia pestis
C.Species: Oz-Nov-2001 #sequence_revision 02-Nov-2001 #sequence_revision 02-Nov-2001 #sequence_revision 02-Nov-2001 #sequence_revision 02-Nov-2001
C.Accession: AE0169
B.Parkhill, J.; Wren, B.W.; Thomson, N.R.; Titball, R.W.; Holden, M.T.G.; Prentice, M.B. Il, M.; Rutherford, K.; Simmonds, M.; Skelton, J.; Stevens, R.M.; Davis, P.; Dougan, G.; Nature 413, 523-527, 2001
A.Fitle: Genome sequence of Yersinia pestis, the causative agent of plague.
A.Reference number: AB0001; MUID:21470413; PMID:11586360
A.Status: preliminary
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          C,Accession: T40491
R;Wood, V.; Rajandream, M.A.; Barrell, B.G.; Lauber, J.; Hilbert, H.; Duesterhoeft, A. Brwood, V.; Rajandream, M.A.; Barrell, B.G.; Lauber, J.; Hilbert, H.; Duesterhoeft, A. Submitted to the EMBL Data Library, February 1998
A;Accession: T40491
A;
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C;Genetics:
A;Gene: YPO1387
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Species: Schizosaccharomyces pombe
Date: 03-Dec-1999 #sequence_revision 03-Dec-1999 #text_change 03-Dec-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 hypothetical protein SPBC4C3.06 - fission yeast (Schizosaccharomyces pombe)
                                                                                                                                                                                                                                                                                                                                                                                                                                   Gaps
A; Residues: 1-1162 <SUT>
A; Cross-references: GB:M21515; NID:9808698; PIDN:AAA66578.1; PID:9331187
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
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                                                                                                                                                                                                                                                                                                                                         Score 52; DB 1; Length 1162;
Pred. No. 39;
4; Mismatches 11; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 818;
                                                                                                                                                                                                                                                                                                                                                                                                                        11; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  2; Length 364;
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Pred. No. 15;
5; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     8; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 4 QVSSPIYD--INYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          31.4%; Score 51.5; 40.0%; Pred. No. 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 472 IDIFVVQSEYGLNYYKVNPCEDVN 495
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1 MDYQVSSPIYDINYYTSEPCOKIN 24
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12; Conservative

Best Local Similarity Matches 12; Conserv

Query Match

66

δ 엄 RESULT 12

5;

31.1%;

Query Match
Best Local Similarity 50.0.

1-364 <KUR>

A;Residues:

A; Molecule type: DNA

1 MDYQVSSPIYDINYYTSE 18

à

A; Accession: S14598
A; Status: preliminary
A; Molecule type: genomic RNA
A; Residues: 1-50 < CAV.>
A; Cross-references: BMBL:X58065; NID:959025; PIDN:CAA41096.1; PID:959026
C; Superfamily: coronavirus E2 glycoprotein

Query Match 31.1%; Score 51; DB 2; Length 520; Best Local Similarity 46.7%; Pred. No. 22; Matches 7; Conservative 3; Mismatches 5; Indels

0;

0; Gaps

10 YDINYYTSEPCQKIN 24 | : | | | | : : | 464 YGINYYKVNPCEDVN 478

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Search completed: July 29, 2004, 13:30:11 Job time : 17 secs

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GenCore version 5.1.6
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- protein search, using sw model OM protein July 29, 2004, 13:24:41 ; Search time 14 Seconds (without alignments) 115.298 Million cell updates/sec Run on:

Title: Perfect score: Sequence:

US-09-852-238A-5 164 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31

Scoring table:

BLOSUM62 Gapop 10.0 , Gapext 0.5

141681 Total number of hits satisfying chosen parameters:

141681 seqs, 52070155 residues

Searched:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 08
Maximum Match 1008
Listing first 45 summaries

SwissProt\_42:\* Database : Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

	Description		בי					P56441 papio hamad			~ 1	m	Φ.		0			_		P51683 mus musculu								Q8sag3 vitis vinif					0206 avian	P30207 avian infec
SUMMAKIES	ID	CKRS HIMAN	Try Carlo	ALCED A	CKR5 CERTO		CKR5 MACMU	CKR5_PAPHA	CKR5_PONPY	CKR5 PYGBI	CKR5 PYGNE	CKR5_TRAFR	CKR5_TRAPH	CKR5_GORGO	CKR5_HYLML	CKR5_CERAE	CKRS_HYLLE	CKR5_MOUSE	CKR5_RAT	CKR2_MOUSE	CKR2_RAT	ADF2_ARATH	ADF4_ARATH	UPP_LACLA	UPP_LACLC	ADF1_PETHY	ADF6 ARATH			VGL2_IBVK	LMB1_DROME	ADF1_ARATH	VGL2_IBVU1	VGL2_IBVU2
	DB	-	۱,-	۱.			-1	Н	Н	Н	-	-	<b>,</b>	⊣	,-	Н	⊣	Н	1	Н	-	Н	H	-	Н	Н	1	-	Н	Н	Н	Н	Н	Н
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	* Query Match					63.6					93.3	93.3	93.3	90.9	9.68	88.4	87.8	64.0	64.0	33.8	33.8	33.2	33.2	32.9	32.9	32.6	32.6	32.3	32.3	31.7	31.7	31.1	31.1	31.1
	Score	164	0 4 5	100	154	154	154	154	154	153	153	153	153	149	147	145	144	105	105	55.5	55.5	54.5	54.5	54	54	53.5	53.5	53	53	52	52	51	51	51
	Result No.			4 0	n 4	יני	ω.	7	80	Q	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33

P30208 avian infec P17662 avian infec	P12722 avian infec P11223 avian infec								
VGL2_IBVU3 VGL2_IBVD3	VGL2_IBVD2 VGL2_IBVB	VGL2_IBV6	OPPF STRP8	OPPF STRPY	LEU3_BACTN	CKR2 MACMU	CKR2 HUMAN	ADF BRANA	VGLL_HSVSA
цц		Н	Н	Н	Н	Н	н	П	ч
520	1154	1163	307	307	353	360	374	126	141
31.1	31.1	31.1	29.9	29.9	29.9	29.9	29.9	29.6	29.6
51	51	51	49	49	49	49	49	48.5	48.5
34 35	36	38	33	40	41	42	43	44	45

## ALIGNMENTS

Ho D.D.;

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EMBL, AF011535; AAB65735.1; BEBL, AF011535; AAB65736.1; BEBL, AF011537; AAB65737.1; BEBL, AF031237; AAB6737.1; BEBL, AF032337; AAB94735.1; BEBL, AF052539; AAD18131.1; Genew, HGNC:1606; CCR5.
               X91492; CAA62796.1; -. U54994; AAC56598.1; -. U57840; AAB17071.1; -. U95626; AAB57793.1; -. U83326; AAC51797.1; -.
                                                                                                         AF011500; AAB65700.1; -.
AF011501; AAB65701.1; -.
AF011502; AAB65702.1; -.
AF011503; AAB65703.1; -.
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                                                                                                                                                                                   AF011505; AAB65705.1;
                                                                                                                                                                                                  AF011506; AAB65706.1;
AF011507; AAB65707.1;
                                                                                                                                                                                                                                                                                                                                              AF011514; AAB65714.1;
AF011515; AAB65715.1;
AF011516; AAB65716.1;
AF011517; AAB65717.1;
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AF011534; AAB65734.1;
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AF011527; AAB65727.1;
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AF011531; AAB65731.1;
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                                                                                                                                                                                                                                       AF011508; AAB65708.1
                                                                                                                                                                                                                                                      AF011509; AAB65709.1
AF011510; AAB65710.1
                                                                                                                                                                                                                                                                                                             AF011512; AAB65712.1
AF011513; AAB65713.1
                                                                                                                                                                                                                                                                                                                                                                                                                                                           AF011520; AAB65720.1
AF011521; AAB65721.1
                                                                                                                                                                                                                                                                                            AF011511; AAB65711.1
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AF011523; AAB65723.1
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha,
                                                   В,
                                                                                                                                                    MEDLINE=98049523; PubMed=9388201; Mummidi S., Ahuja S.K.; Mummidi S., Ahuja S.S., McDaniel B.L., Ahuja S.K.; Mummidi S., Ahuja S.S., McDaniel B.L., Ahuja S.K.; The human CC chemokine receptor 5 (CCR5) gene. Multiple transcripts with 5'-end heterogeneity, dual promoter usage, and evidence for Jolymorphisms within the regulatory regions and noncoding exons."; J. Biol. Chem. 272:30662-30671(1997).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MIP-1-beta and rates and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or differentiation. Acts as co-eceptor with CD4 for primary non-syncytium-inducing strains (NSI) (macrophage-tropic) of HIV-1 virus. It promotes Env-mediated fusion of the virus. SUBCELLULAR LOCATION: Integral membrane protein.

TISSUE SPECIFICITY: Found in promyelocytic cells.

PTM: Sulfation contributes to the efficiency of HIV-1 entry.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     MEDLINE=99189752; PubMed=10089882; Farzan M., Mirzabekov T., Kolchinsky P., Wyatt R., Cayabyab M., Gerard N.P., Gerard C., Sodroski J., Choe H.; "Tyrosine sulfation of the amino terminus of CCR5 facilitates HIV-1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       glycosylation. SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
         SEQUENCE FROM N.A., AND POLYMORPHISMS.
MEDLINE-98022612; PubMed-9359654;
Zhang L., Carruthers C.D., He T., Huang Y., Cao Y., Wang G., Hahn
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Paxton W.A.; "HIV-1 entry into CD4+ cells is mediated by the chemokine receptor
                                                                                "HIV type 1 subtypes, coreceptor usage, and CCR5 polymorphism."; AIDS Res. Hum. Retroviruses 13:1357-1366(1997).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CHARACTERIZATION OF 1TS HIV-1 RECEPTOR FUNCTION.
MEDLINE=96260018; PubMed=8649512;
Dragic T., Litwin V., Allaway G.P., Martin S.R., Huang Y.,
Nagashima K.A., Cayanan C., Maddon P.J., Koup R.A., Moore J.P.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     MEDLINE=96260017; PubMed=8649511;
Deng H., Liu R., Ellmeler W., Choe S., Unutmaz D., Burkhart M., di Marzio P., Marmon S., Sutton R.E., Hill C.M., Davis C.B., Peiper S.C., Schall T.J., Littman D.R., Landau N.R.;
"Identification of a major co-receptor for primary isolates of
                                                                                                                                                                                                                                                                                                                                                                                            Kopatz S.A., Aronstam R.S., Sharma S.V.; "cDNA clones of human proteins involved in signal transduction sequenced by the Guthrie cDNA resource center (www.cdna.org)."; Submitted (JAN-2003) to the EMBL/GenBank/DBJ databases.
                                                                                                                                                                                                                                                                                 SEQUENCE FROM N.A., AND VARIANT ARG-178.
Magierowska M., Barre-Sinoussi F., Issafras H., Theodorou
                                                                                                                                                                                                                                                                                                                                          Submitted (MAR-1998) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CHARACTERIZATION OF ITS HIV-1 RECEPTOR FUNCTION.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  HIV-1.";
Nature 381:661-666(1996).
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Cell 96:667-676(1999)
                                                                                                                                       SEQUENCE FROM N.A.
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CC-CKR-5

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MIM; 601373; -.

R GO; GO:000576B; C:endosome; TAS.

R GO; GO:00058B7; C:integral to plasma membrane; TAS.

R GO; GO:0016493; F:C-C chemokine receptor activity; NAS.

R GO; GO:0016493; F:C-C chemokine receptor activity; NAS.

R GO; GO:00165026; F:coreceptor activity; TAS.

R GO; GO:0005935; P:cellular defense response; TAS.

R GO; GO:0006935; P:chemotaxis; TAS.

R GO; GO:0007204; P:cytosolic calcium ion concentration elevation; TAS.

R GO; GO:0007204; P:cytosolic calcium ion concentration elevation; TAS.

R GO; GO:0007205; P:inflammatory response; TAS.

R GO; GO:0007203; P:phosphatidylinositol-4,5-bisphosphate hydro. .; TAS.

InterPro; IRR000276; GPCR R.Rhodpsn.

R Pfam; PP00001; 7tm 1.1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  PROSITE; PS00237; G_PROTEIN_RECEP_F1_1; 1.
PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
G_Protein coupled receptor; Transmembrane; Glycoprotein; Sulfation; Polymorphism.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1 (POTENTIAL).
CYTOPIASMIC (POTENTIAL).
2 (POTENTIAL).
EXTRACELLUIAR (POTENTIAL).
3 (POTENTIAL).
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В.
                                                                 Gaps
                                                                                                                                                                                                                                                           CKRS_PANTR STANDARD; PRT; 352 AA.
PS6440; 002778;
15-UTL-1998 (Rel. 36, Last sequence update)
15-UTL-1998 (Rel. 36, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CS) (CCRS).
CCRS OR CMKBRS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       -:- SUBCELLULAR LOCATION: Integral membrane protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Zacharova V., Zachar V., Goustin A.S.; "Sequence of chemokine receptor gene CCR5 in chimpanzees, a natural
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SEQUENCE FROM N.A.
MEDIINE=97268687; PubMed=9108095;
Edinger A.L., Amedee A., Miller K., Doranz B.J., Endres M.,
Sharron M., Samson M., Lu Z.H., Clements J.E., Murphey-Corb M.,
Peiper S.C., Parmentier M., Broder C.C., Doms R.W.;
"Differential utilization of CCR5 by macrophage and T cell tropic
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Zhang L., Carruthers C.D., He T., Huang Y., Cao Y., Wang G., Hahn
                                                                                                                                                                                                                                                                                                                                                                                                                                     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Pan.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  "HIV type 1 subtypes, coreceptor usage, and CCR5 polymorphism."; AIDS Res. Hum. Retroviruses 13:1357-1366(1997).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MEDLINE=98090115; PubMed=9430250; Pretet J.-G., Butor C.; Pretet J.-L., Zerbib A., dirard M., Guillet J.-G., Butor C.; Chimpanzee CXCR4 and CCR5 act as coreceptors for HIV type 1."; AIDS Res. Hum. Retroviruses 13:1583-1587(1997).
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                      Length 352;
                                                             0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SEQUENCE FROM N.A.
Zimmerman P.A., Buckler-White A., Alkhatib G.;
Submitted (MAR-1997) to the EMBL/GenBank/DDBJ databases.
                  ; Score 164; DB 1;
; Pred. No. 5.8e-16;
0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        simian immunodeficiency virus strains.";
Proc. Natl. Acad. Sci. U.S.A. 94:4005-4010(1997)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     HIV type 1 host.";
AIDS Res. Hum. Retroviruses 13:1159-1161(1997).
                                                                                                                                   1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                        1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MEDLINE=97426118; PubMed=928282;
                      100.08;
                                                                                                                                                                                                                                                                                                                                                                                                                 Pan troglodytes (Chimpanzee).
                                                               31; Conservative
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                                           Similarity
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SEQUENCE FROM N.A.
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Best Local &
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use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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15-MAR-2004 (Rel. 43, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR-5) (CCRS).
                                                                                                                                                                                                                                       PFam; PF00001; /rum 1; PF000027; GPCRHDODESN.
PRINTS; PR00237; GPCRHDODESN.
PROSITE; PS00262; GPROTEIN_RECEP_F1_2; 1.
PROSITE; PS50262; GPROTEIN_RECEP_F1_2; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
30 EXTRACELLULAR; (POTENTIAL).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Muller-Trutwin M.C., Corbet S., Hansen J., Georges-Courbot M.-C., Diop O., Rigoulet J., Barre-Sinoussi F., Fomsgaard A.; "Mutations in CCR5-coding sequences are not associated with SIV carrier status in African nonhuman primates.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Cercopithecus pygerythrus (Vervet monkey).
Wakaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae;
Cercopithecinae; Cercopithecus.
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N REF. 1).
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EXTRACELLULAR (POTENTIAL).
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SULFATION (BY SIMILARITY)
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Pred. No. 3e-15;
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MEDLINE=99335215; Pubmed=10408730;
                                                                                                                                                                                EMBL; U89797; AAC03717.1; -.
EMBL; AF177894; AAK43377.1; -
InterPro; IPR000276; GPCR_Rhodpsn.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     15-MAR-2004 (Rel. 43, Created)
15-MAR-2004 (Rel. 43, Last sequ
15-MAR-2004 (Rel. 43, Last anno
                                                                                                                            EMBL; AF011542; AAB65742.1; -. EMBL; U97666; AAC51670.1; -.
                                                                                                                                                               EMBL; AF011540; AAB65740.1; -.
                                                                                            EMBL; AF005663; AAB62557.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                40539 MW;
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96.8%;
                                                                                                              U94329; AAB58446.1;
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352 AA;
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CKR5_CERPY
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"Primary SIVsm isolates use the CCR5 coreceptor from sooty mangabeys
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                                                                                                 This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch)
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AIDS Res. Hum. Retroviruses 15:931-939(1999).
-!-FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gaps
                                                                                                                                                                              EMBL; AF0352222, AAD44015.1; -.
Interpro; IPR000276, GPCR_Rhodpsn.
PRINTS; PR00037; GFCRHODOPSN.
PROSITE; PS00237; GFROTEIN RECEP F1 1; 1.
PROSITE; PS500262; GFROTEIN RECEP F1 2; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              CKRS_CERTO STANDARD, PRT; 352 AA. 062745; 062746; 15-DRC-1938 (Rel. 37, Created) 15-DEC-1938 (Rel. 37, Created) 15-DEC-1938 (Rel. 37, Last sequence update) 16-OCT-2001 (Rel. 40, Last amoration update) 16-OCT-2001 (Rel. 40, Last amoration update) 16-CC Chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCRS).
                                                     differentiation.
-!- SUBCELLULAR LOCATION: Integral membrane protein.
-!- SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Cercocebus torquatus atys (Red-crowned mangabey) (Sooty mangabey).
Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 352;
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                                                                                                                                                                                                                                                                                                                                                                                                                                          BY SIMILARITY.
SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
                                                                                                                                                                                                                                                                                                                                                               EXTRACELLULAR (POTENTIAL)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      EF17D67CBCCC3DB0 CRC64;
                                                                                                                                                                                                                                                                                                                  EXTRACELLULAR (POTENTIAL)
                                                                                                                                                                                                                                                                                                                              3 (POTENTIAL).
CYTOPLASMIC (POTENTIAL).
                                                                                                                                                                                                                                                                                                                                                                        5 (POTENTIAL).
CYTOPLASMIC (POTENTIAL).
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2 (POTENTIAL).
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                                                                                                                                                                                                                                                                                                                                                   4 (POTENTIAL).
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                                                                                                                                                                                                                                                                                                                                                                                                (POTENTIAL)
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                                                                                                                                                                                                                                                                                                                                                                                                                      7 (POTENTIAL)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQUENCE FROM N.A.
STRAIN=Isolate 079, 085, 087, and 089;
MEDLINE=98321155; PubMed=9656999;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      40558 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           93.9%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Local Similarity 93.5
                                                                                                                                                                                                                                                                                            68
89
102
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235
260
277
277
301
352
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naturally infected in west Africa: a comparison of coreceptor usage of primary SIV8m, HIV-2, and SIVmac."; virology 246:113-124(1998).
-!- FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or differentiation.
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EMBL, AF051904; AAC39833.1; --
EMBL, AF051905; AAC39833.1; --
INCEPPRO, IRROMODES, APC39833.1; --
INCEPPRO, IRROMODES, APC3983.1; --
PROMODI, 7tm 1; 1.
PRINTS; PR00237; GPCRRHODOPSN.
PROSITE, PS00237; GPCRRHODOPSN.
PROSITE, PS00237; GPROTEIN RECEP_F1 1; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
DOMAIN
                                                                                                                                                                                 SUBCELLULAR LOCATION: Integral membrane protein. SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
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Y -> D (IN ISOLATE 087).

Y -> G (IN ISOLATE 087).

M -> K (IN ISOLATE 087).

L -> V (IN ISOLATE 089).

V -> G (IN ISOLATE 089).

Y -> G (IN ISOLATE 089).

Y -> L (IN ISOLATE 089).

Y -> I (IN ISOLATE 089).

Y -> I (IN ISOLATE 079).

W, 20A196E2D47E49CA CRC64;
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CYTOPLASMIC (POTENTIAL).
4 (POTENTIAL).
EXTRACELULAR (POTENTIAL).
5 (POTENTIAL).
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SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
D -> E (IN ISOLATE 087).
Y -> D (IN ISOLATE 087).
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EXTRACELLULAR (POTENTIAL)
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CYTOPLASMIC (POTENTIAL).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  1; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                   EMBL; AF051902; AAC39830.1; -. EMBL; AF051903; AAC39831.1; -.
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278
302
101
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Q95NCS;
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SEQUENCE FROM N.A.
SPECIES=M.mulatta; STRAIN=Indian macaque;
SPECIES=M.mulatta; PubMed=9060623;
Chen Z., Zhou P., Ho D.D., Landau N.R., Marx P.A.;
"Genetically divergent strains of simian immunodeficiency virus use
"Genetically divergent for entry.";
J. Virol. 71:2706-2714(1997).
 01-NOV-1997 (Rel. 35, Last sequence update)
28-PEB-2003 (Rel. 41, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR-5) (CCRS)
CCRS OR CMKBRS.
                                                                         Macaca fascicularis (Crab eating macaque) (Cynomolgus monkey), and
Macaca nemestrina (Pig-tailed macaque).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MEDIINE=21354176; PubMed=11461684;
Margulies B.J., Hauer D.A., Clements J.E.;
"Identification and comparison of eleven rhesus macaque chemokine
                                                                                                                                                                                                           MEDLINE=97184592; PubMed=9032394;
Marcon L., Choe H., Martin K.A., Farzan M., Ponath P.D., Wu L., Newman W., Gerard N., Gerard C., Sodroski J.;
"Utilization of C-C chemokine receptor 5 by the envelope glycoproteins of a pathogenic simian immunodeficiency virus, SIVmac239.";
                                                                                                        Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Eutele
Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae,
                                                           Macaca mulatta (Rhesus macaque),
                                                                                                                                                                                                                                                                                                     Virol. 71:2522-2527(1997).
                                                                                                                                      Cercopithecinae, Macaca.
NCBI_TaxID=9544, 9541, 9545;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SPECIES=M.mulatta;
                                                                                                                                                                                                SPECIES=M.mulatta;
                                                                                                                                                                                SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                       FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    receptors."
                                                                                                                                                                                                                                                                                                                                                                                                                                                       SEQUENCE
    This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the BMBL outstation - the Buropean Bioinformatics Institute of There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@ib-sib.ch).
                                                                                                                                                             MEDLINE=99416438; PubMed=10486970; Zhang Y.-W., Ryder O.A., Zhang Y.-P.; Zhang Y.-W., Ryder O.A., Zhang Y.-P.; Sequence evolution of the CCR5 chemokine receptor gene in primates."; Mol. Biol. Evol. 16:1145-1154(1999).
-!- FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
15-MAR-2004 (Rel. 43, Created)
15-MAR-2004 (Rel. 43, Last sequence update)
15-MAR-2004 (Rel. 43, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR-5)
                                                                                                                                                                                                                                                                                                                  -!- SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                InterPro; IPR000276; GPCR_Rhodpsn.
Pfam; PF00001; 7tm 1; 1.
PRINTS; PR00237; GPCRRHODDPSN.
PROSTIE; PS00237; G PROTEIN RECEP F1 1; 1.
PROSITE; PS50262; G PROTEIN RECEP F1 2; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
                                                                         Hylobates syndactylus (Siamang) (Symphalangus syndactylus).
Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hylobatidae, Hylobates.
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EXTRACELLULAR (POTENTIAL)
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EXTRACELLULAR (POTENTIAL).
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SULFATION (BY SIMILARITY)
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CYTOPLASMIC (POTENTIAL).
4 (POTENTIAL).
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CYTOPLASMIC (POTENTIAL).
                                                                                                                                                                                                                                                                                                  SUBCELLULAR LOCATION: Integral membrane protein.
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301
352
178
                                                                                                                                                                                                                                                                                      differentiation.
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352 AA;
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nes 29; Conserv
                                                                                                                                                  SEQUENCE FROM N.A.
                                                                                                                     NCBI_TaxID=9590;
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125
142
                                                            CCR5 OR CMKBR5.
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                                                                                                                    SPECIBE=M. m.n.n. A. fascicularis, and M.nemestrina, processes from N.n. Amedee A., Miller K., Doranz B.J., Endres M., Bidnger A.L., Amedee A., Miller K., Doranz B.J., Endres M., Sharron M., Samson M., Lu Z.H., Clements J.E., Murphey-Corb M., Peiper S.C., Parmentier M., Broder C.C., Doms R.W.; "Differential utilization of CCR5 by macrophage and T cell tropic simian immunodeficiency virus strains.", Proc. Natl. Acad. Sci. U.S.A. 94.4005-4010(1997).

-I. FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RAWIES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or differentiation.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            -!- SUBCELLULAR LOCATION: Integral membrane protein.
-!- SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
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EMBL, AP005661; AAB62554.1; -.
EMBL, AP005662; AAB62555.1; -.
EMBL, AP005662; AAB62556.1; -.
InterPro; IRR000276; GPCR_Rhodpsn.
Pfan, PF00017, 7m. 1; 1.
Pfan, PF00017, 7m. 1; 1.
PROSITE; PS00237; GPRRHODOPSN.
AIDS Res. Hum. Retroviruses 17:981-986(2001)
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Gaps

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Indels

1; Mismatches

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352 AA.

PRT;

STANDARD;

CKR5 MACMU

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CKR5\_MACMU

RESULT 6

P79436; 002746; 01-NOV-1997 (Rel. 35, Created)

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              15-JUL-1998 (Rel. 36, Created)
15-JUL-1998 (Rel. 36, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
16-OCT-Comemokine receptor type 5 (C-C Chemokine receptor type 5 (C-C CRR-5) (CC-CRR-5) (CCRS).
CCRS OR CMRBRS.
Papio hamadryas (Hamadryas baboon), and
G_PROTEIN RECEP F1 2; 1. receptor; Transmembrane; Glycoprotein; Sulfation. 30 EXTRACELLULAR (POTENTIAL).
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MEDLINE=97268687; PubWed=9108095;
Edinger A.L., Amedee A., Miller K., Doranz B.J., Endres M.,
Sharron M., Samson M., Lu Z.H., Clements J.E., Murphey-Corb M.,
Peiper S.C., Parmentier M., Broder C.C., Doms R.W.;
"Differential utilization of CCR5 by macrophage and T cell tropic
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae,
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Benton P.A., Timanus D.K., Shearer M.H., Lee D.R., Kennedy R.C.;
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SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
SULFATION (BY SIMILARITY).
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EXTRACELLULAR (POTENTIAL)
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4 (POTENTIAL).
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CYTOPLASMIC (POTENTIAL).
                                                                          1 (POTENTIAL).
CYTOPLASMIC (POTENTIAL).
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Proc. Natl. Acad. Sci. U.S.A. 94:4005-4010(1997)
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-> M (IN REF. 3)
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AIDS Res. Hum. Retroviruses 15:479-483(1999)
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29; Conservative
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NCBI_TaxID=9557, 9555;
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352 AA;
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Best Local Similarity
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SEQUENCE FROM N.A.
    PS50262;
                                 G-protein coupled
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differentiation.
-!-SUBCELLUIAR LOCATION: Integral membrane protein.
-!-SUBCELLUIAR LOCATION: Integral membrane protein.
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             -!- FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gaps
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30-MX-2000 (Rel. 39, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
28-FEB-2003 (Rel. 41, Last annotation update)
CCC chemokine receptor type 5 (C-C CKR-5) (CCRS).
CCR5 OR CMKBR5.
Pongo pygmaeus (Orangutan).
Ebukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Pongo.
                                                                                                                                                                                                                                                                                                                                                                         PROSITE; PS00237; G_PROTEIN_RECEP_F1_1; 1.
PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
G_PTOCLEIN COUPLED ** CCEPTOR** Transmembrane; Glycoprotein; Sulfation.
DOMAIN 1 30 EXTRACELLULAR (POTENTIAL).
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Submitted (NOV-1997) to the EMBL/GenBank/DDBJ databases.
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S (POTENTIAL).
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(BY SIMILARITY)
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5E1504A9BA1FE8B2 CRC64;
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EXTRACELLUIAR (POTENTIAL)
3 (POTENTIAL).
CYTOPLASMIC (POTENTIAL).
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SULFATION (BY SIMILARITY)
SULFATION (BY SIMILARITY)
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CYTOPLASMIC (POTENTIAL).
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EMBL; AF105287; AAD20556.1; -.
EMBL; AF105288; AAD20557.1; -.
EMBL; AF105289; AAD20558.1; -.
EMBL; AF105290; AAD20559.1; -.
EMBL; AF02452; AAC63830.1; -.
InterPro; IRROGO276; GPCR_Rhodpsn.
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89
102
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
      Pygathrix.
NCBI_TaxID=61621;
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ID CKRS_PYGNE
AC 097882;
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                                                                                                                                                                                                             ong as its content is in no way toved. Usage by and for commercial (See http://www.isb-sib.ch/announce/
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                                           Zhang Y. W., Ryder O.A., Zhang Y.-P.; "Sequence evolution of the CCR5 chemokine receptor gene in primates."; Mol. Biol. Evol. 16:1145-1154(1999)
                                                                            FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR5).
CCR5 OR CMKRRS.
Pygathrix bieti (Black snub-nosed monkey) (Rhinopithecus bieti).
Eukaryota, Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae; Colobinae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                             InterPro; 1PRUDUZZ,, 1 ...
PERM: PRODOS1; 7H 1; 1.
PROSITE; PRO0237; GPRAPIN RECEP F1 1; 1.
PROSITE; PS50262; G PROTEIN RECEP F1 2; 1.
PROSITE; PS50262; G PROTEIN RECEP F1 2; 1.
G-protein cupled receptor; Transmembrane, Glycoprotein; Sulfation.
31 30 EXTRACELUIAR (POTENTIAL).
31 58 I (POTENTIAL).
                                                                                                                                      -!- SUBCELLULAR LOCATION: Integral membrane protein.
-!- SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
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CYTOPLASMIC (POTENTIAL).
4 (POTENTIAL).
EXTRACELLULAR (POTENTIAL)
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EXTRACELLULAR (POTENTIAL)
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SULFATION (BY SIMILARITY)
SULFATION (BY SIMILARITY)
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EXTRACELLULAR (POTENTIAL)
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Pred. No. 1.5e-14;
                                                                                                                                                                                                                   modified and this statement is not removed.
entities requires a license agreement (See hor send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1 MDYQVSSPTYDIDYYTSBPCQKINVKQIAAR 31
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                     SEQUENCE FROM N.A.
MEDLINE=99416438; PubMed=10486970;
                                                                                                                                                                                                                                                                 EMBL; AF075446; AAD19858.1; -.
InterPro; IPR000276; GPCR_Rhodpsn.
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l Similarity 93.5%;
29; Conservative
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                                                                                                                            differentiation.
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 NCBI_TaxID=9600;
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097880;
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                        MEDLINE=99416438; PubMed=10486970; Zhang Y.-W.; Kyder O.A., Zhang Y.-P.; "Sequence evolution of the CCR5 chemokine receptor gene in primates."; Mol. Biol. Evol. 16:1145-1154(1999)
                                                                                                                         -!- FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
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                                                                                                                                                                                                                                                          -!- SUBCELLULAR LOCATION: Integral membrane protein.
-!- SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  InterPro; IPR000276; GPCR Rhodpsn.
Pfam; PF00001; 7tm 1; 1.
PRINTS; PR000237; GPCRRHODDPSN.
PROSITE; PS00237; G PROTEIN RECEP F1 1; 1.
PROSITE; PS50262; G PROTEIN RECEP F2; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
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30-MAY-2000 (Rel. 39, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKS).
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EXTRACELLULAR (POTENTIAL)
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SULFATION (BY SIMILARITY)
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Pred. No. 2.1e-14;
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SEQUENCE FROM N.A.
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                                                                              SEQUENCE FROM N.A.

MEDLINES-99416438; PubMed=10486970;

Zhang Y.-W., Ryder O.A., Zhang Y.-P.;

Zhang Y.-W., Ryder O.A., Zhang Y.-P.;

"Sequence evolution of the CCRS chemokine receptor gene in primates.";

Mol. Biol. Evol. 16:1145-1154(1999).

II-FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha,

MIP-1-beta and RANTES and subsequently transduces a signal by

increasing the intracellular calcium ions level. May play a role

in the control of granulocytic lineage proliferation or

differentiation.
Chordata, Craniata, Vertebrata, Buteleostomi;
Primates, Catarrhini, Cercopithecidae, Colobinae;
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30-MAY-2000 (Rel. 39, Last sequence update)
28-FBB-2003 (Rel. 41, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKF) (CCRS).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PROSITE; PS00237; G_PROTEIN_RECEP_FI_1; 1.
PROSITE; PS50262; G_PROTEIN_RECEP_FI_2; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
DOMAIN 1 30 EXTRACELLULAR (POTENTIAL).
                                                                                                                                                                                                                                                           -!- SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
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EXTRACELLULAR (POTENTIAL)
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EXTRACELLULAR (POTENTIAL)
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CYTOPLASMIC (POTENTIAL).
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CYTOPLASMIC (POTENTIAL)
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2.1e-14;
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InterPro; PR000275; GPCR_Rhodpsn.
Pfam. PF00001; 7tm 1; 1.
PRINTS; PR00237; GPCRRHODOPSN.
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Matches 28; Conservative
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15
                     Mammalia; Eutheria;
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                                                  NCBI_TaxID=54180;
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278
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CKR5_TRAPH
ID CKR5_TRAPH
097879;
^-MAY-20'
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                                                                                                                                                                                                                                                                        This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstationthe European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (see http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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                                                                                                  Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae; Colobinae;
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30-MAY-2000 (Rel. 39, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CS) (CCRS).
CCRS OR CMKBRS.
                                                                                                                                                                                                                                                                                                                                                                                            EMBL, AF075448; AAD19860.1; -.

InterPro; IPR000276; GPCR_Rhodpsn.

Pfam; PF00001; 7tm 1; 1.

PRINTS; PR00237; GPCRHODOPSN.

PROSITE; PS00237; GPROTEIN RECEP_F1_1; 1.

PROSITE; PS50262; GPROTEIN_RECEP_F1_2; 1.

G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.

G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
                                                                                                                                                                                                                                SUBCELLULAR LOCATION: Integral membrane protein. SIMILARITY: Belongs to family 1 of G-protein coupled receptors
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CYTOPLASMIC (POTENTIAL).
6 (POTENTIAL).
EXTRACELLULAR (POTENTIAL)
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FE4F9D98D3B3E861 CRC64;
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EXTRACELLULAR (POTENTIAL)
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CYTOPLASMIC (POTENTIAL).
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Pred. No. 2.1e-14;
2; Mismatches 1;
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                                                                      SEQUENCE FROM N.A.
MEDLINE=99416438; PubMed=10486970;
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Matches 28; Conservative
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NCBI_TaxID=54133;
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CKR5 HYLML
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ò
                                                                                                                  MEDLINE=99416438; PubMed=10486970; Zhang Y.-P.; Zhang Y.-W., Ryder O.A., Zhang Y.-P.; Full Shang Y.-P.; Zhang Y.-P.; Z
                Eukaryota; Metažoa; Chordatá; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae; Colobinae;
                                                                                                                                                                                                                                                                                                                                                                                                                                         or send an.

EMBL, AF075443; AAD19855.1; -

EMBL, AF075443; AAD19855.1; -

InterPro; IPR000276; GPCR.Rhodpsn.

PRIMTS; PR000237; GPCRRHODOSN.

PROSITE; PS00237; GPCRRHODOSN.

PROSITE; PS00262; G—PROTEIN_RECEP_F1_2; 1.

G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.

G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
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15-JUL-1998 (Rel. 36, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR-5).
                                                                                                                                                                                                                                                                               SUBCELLULAR LOCATION: Integral membrane protein. SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
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(BY SIMILARITY).
(BY SIMILARITY).
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2 (POTENTIAL).
EXTRACELLULAR (POTENTIAL).
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EXTRACELLULAR (POTENTIAL)
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CYTOPLASMIC (POTENTIAL).
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Trachypithecus phayrei (Phayre's leaf monkey)
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SULFATION (BY
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352 AA;
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                                                                                                       SEQUENCE FROM N.A.
                                                                     NCBI_TaxID=61618;
                                                    Trachypithecus
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P56439;
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CKR5 GORGO
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                                                                                                                                              MEDLINE=97268687; PubMed=9108095; Bdinger A.L., Amedee A., Miller K., Doranz B.J., Endres M., Sharron M., Samson M., Lu Z.H., Clements J.E., Murphey-Corb M., Peiper S.C., Parmentier M., Broder C.C., Dons R.W., "Differential utilization of CCRS by macrophage and T cell tropic simian immunodeficiency virus strains."; Proc. Natl. Acad. Sci. U.S.A. 94:4005-4010(1997).

-!- FUNCIION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RAWTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
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InterPro; IPR00276; GPCR_Rhodpsn.
Prom; PR00021; 7tm 1, 1.
PROSTIE; PR00237; GPCRHUDOPSN.
PROSTIE; PS00262; GPROTEIN RECEP_F1 1; 1.
PROSTIE; PS0262; GPROTEIN TRECEP_F1 2; 1.
DOMAIN : 30 EXTRACELLULAR (POTENTAL).
                                                                                                                                                                                                                                                                                                                                                                                                                                                           -!- SUBCELLULAR LOCATION: Integral membrane protein.
Gorilla gorilla gorilla (Lowland gorilla).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Gorilla.
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EXTRACELLULAR (POTENTIAL)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       3 (POTENTIAL).
CYTOPLASMIC (POTENTIAL).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         352 AA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1 MDYQVSSPTYDIDYYTSEPCOKTNVKQIAAR 31
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095NCO;
15-MAR-2004 (Rel. 43, Created)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               40515 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  90.06
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                                                                                                                                                                                                                                                                                                                                                                                                                                        differentiation.
                                                                                                                              SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               352 AA;
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                                                                        NCBI_TaxID=9595;
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Submitted (JUL-1998) to the EMBL/GenBank/DDBJ databases.
-!- FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha
MIP-1-beta and RANTES and subsequently transduces a signal by
increasing the intracellular calcium ions level. May play a role
in the control of granulocytic lineage proliferation or
           chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR-5) (CCR5)
                                                                                                                                                    MEDLINE=98001387; PubMed=9343222; Kuhmann S.E., Platt E.J., Kozak S.L., Kabat D.; Kuhmann s.E., Platt E.J., Kozak S.L., Kabat D.; Prolymorphisms in the CCRS genes of African green monkeys and mice implicate specific amino acids in infections by simian and human immunodeficiency viruses.";
                                      Cercopichecus aethiops (Green monkey) (Grivet).
Bukaryota, Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae;
                                                                                                                                                                                                                                                                Murayama Y., Matsunaga S., Inoue-Murayama M.; "cDNA sequence of African green monkey CCR-5 chemokine receptor
 16-OCT-2001 (Rel. 40, Last annotation update)
                                                                                   Cercopithecinae; Cercopithecus.
                                                                                                                                                                                                                          Virol. 71:8642-8656(1997).
                                                                                                                                                                                                             ency viruses.
                                                                                                                                                                                                                                                                                                                                                                                  differentiation.
                                                                                                                                                                                                                                                      SEQUENCE FROM N.A.
                                                                                                                           SEQUENCE FROM N.A.
                                                                                             NCBI TaxID=9534;
                           CMKBR5.
                                                                                                                                        TISSUE=Kidney
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                             CCR5 OR
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
                                                                                                                                      MEDLINE=994164.38; PubMed=10486970; Zhang Y.-P.; Zhang Y.-P., Zhang Y.-P., Zhang Y.-P.; Jane Zhang Y.-P.; Full State of the CRES chemokine receptor gene in primates."; Mol. Biol. Evol. 16:1145-1154(1999).
-: FUNCTION: Receptor for a C-C type chemokine. Binds to MIP-1-alpha, MIP-1-beta and RANTES and subsequently transduces a signal by increasing the intracellular calcium ions level. May play a role in the control of granulocytic lineage proliferation or
                                                                                                                                                                                                                                                                                                                                                                                                                       EMBL, AF177899, AAK43382.1; -.
InterPro, IPR000276; GPCR_Rhodpsn.
Pfam; PF00001; 7tm 1; 1.
PRINTS; PR00237; GPCRRHODOPSN.
PROSITE; PS00237; G PROTEIN RECEP_F1 1; 1.
PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
G_protein coupled receptor; Transmembrane; Glycoprotein; Sulfation.
  15-MAR-2004 (Rel. 43, Last sequence update)
45-MAR-2004 (Rel. 43, Last annotation update)
C-C chemokine receptor type 5 (C-C CKR-5) (CC-CKR-5) (CCR-5)
                                                                                                                                                                                                                                                                      SUBCELLULAR LOCATION: Integral membrane protein. SIMILARITY: Belongs to family 1 of G-protein coupled receptors.
                                                                      Eŭkaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hylobatidae, Hylobates.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 352;
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SIMILARITY)
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EXTRACELLULAR (POTENTIAL)
3 (POTENTIAL).
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CYTOPLASMIC (POTENTIAL).
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Pred. No. 1.5e-13;
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                                                          Hylobates moloch (Silvery gibbon)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              40436 MW;
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ilarity 90.3%;
Conservative
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                                                                                                                                                                                                                                                         differentiation.
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352 AA;
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Best Local Similarity
Matches 28; Conserv
                                                                                                                               SEQUENCE FROM N.A.
                                                                                                  NCBI_TaxID=81572;
                                           CCR5 OR CMKBR5.
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MOD_RES
SEQUENCE
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Binds to MIP-1-alpha,

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        PEGNI, PF00001; 7cm 1; 1. PRINTS; PR00237; GFCREHODOPSN.
PROSITE; PS00237; GFCREHODOPSN.
PROSITE; PS02262; G PROTEIN_RECEP_F1_1; 1.
G-protein coupled receptor; Transmembrane; Glycoprotein; Sulfation;
-!- SUBCELLULAR LOCATION: Integral membrane protein.
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EXTRACELLULAR (POTENTIAL)
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EXTRACELLULAR (POTENTIAL)
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CYTOPLASMIC (POTENTIAL).
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SULFATION (BY S
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CYTOPLASMIC (
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EMBL; AB015944; BAA31328.1; -.
InterPro; IPR000276; GPCR_Rhodpsn.
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1 MDYQVSSPTYDIDYYTSGPCQKINVKQIAAR 31

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1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31

(Rel. 36, Created) (Rel. 36, Last sequence update)

15-JUL-1998 15-JUL-1998 CERAE

CKR5 CER P56493;

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CKR5\_CERAE RESULT 15

352

STANDARD;

0; Gaps Best Local Similarity 90.3%; Pred. No. 2.9e-13; Matches 28; Conservative 1; Mismatches 2; Indels

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Search completed: July 29, 2004, 13:28:57 Job time: 15 secs

(0102N) AMDIA 8609 2141

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July 29, 2004, 13:25:16; Search time 35 Seconds (without alignments) 279.459 Million cell updates/sec
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GenCore version 5.1.6
Copyright (c) 1993 - 2004 Compugen Ltd.
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1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Total number of hits satisfying chosen parameters:
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Maximum Match 100%
Listing first 45 summaries
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1: SP_archea:*
2: SP_bacteria:*
3: SP_fungi:*
4: Sp_human:*
5: Sp_mammal:*
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Gapop 10.0 , Gapext 0.5
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Maximum DB seq length: 200000000
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Perfect score:
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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SUMMARIES			_	_		٥١	_	~	-			_	•		_			٥.
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PRT; 215 AA.  reated) ist sequence update) ist annotation update)  Craniata; Vertebrata; Euteleostomi;  Catarrhini; Hominidae; Homo.  G., Michael N.L.;  tional Characterization of the Complete mokine Receptor CCR-5, a Major Entry Co-  BL/GenBank/DDBJ databases.  membrane; IEA.  ivity; IEA.  we receptor activity; IEA.  upled receptor activity; IEA.  CEP_FI 1; 1.  CEP_FI 1; 1.  3C9146C76BA416F7 CRC64;  Score 164; DB 4; Length 215;  Pred. No. 4.8e-16;
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1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31

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Query Match
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Matches
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AIDS Res. Hum. Retroviruses 0:0-0(1997).
BIME, AF011539; AAB65739.1; -.
GO; GO:0016021; C:integral to membrane; IEA.
GO; GO:0004872; F:receptor activity; IEA.
GO; GO:0001884; F:rhodopsin-like receptor activity; IEA.
GO; GO:0007186; P:G-procein coupled receptor protein signalin. ..; IEA.
Interpro; IPR000276; GPCR_Rhodpsn.
Pfan; PF00001; 7tm 1; 1.
PRINTS; PR00021; 7tm 1; 1.
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В
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Mammalia, Butheria, Primates, Catarrhini, Hominidae, Pan.
NCBI_TaxID=9598;
                                                                                                                                                                                                                                            Miopithecus talapoin (Talapoin) (Cercopithecus talapoin).
Eukaryota, Metazoa, Chordata, Craniata; Vertebrata, Euteleostomi,
Mammailai, Eutheria, Primates, Catarrhini, Cercopithecidae;
Cercopithecinae; Miopithecus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              EMBL; AF177886; AAK43369.1; --
GO; GO:0016021; C:integral to membrane; IEA.
GO; GO:0004872; F:receptor activity; IEA.
GO; GO:0001384; F:rhodopsin-like receptor activity; IEA.
GO; GO:0007186; F:receptor coupled receptor protein signalin.
InterPro; IPR000276; GPCR Rhodpsn.
PF00001; 7tm 1; 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                             phylogeny.";
Submitted (AUG-1999) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     40546 MW; 6464152F3E566AE5 CRC64;
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Last annotation update)
                                                                                                                                                                  01-DEC-2001 (TrEMBLrel. 19, Last sequence update) 01-JUN-2003 (TrEMBLrel. 24, Last annotation update) C-C chemokine receptor 5.
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4.4e-15;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            97.0%; Score 159; DB 96.8%; Pred. No. 4.4e-
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                                                                                                       352 AA
1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
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PROSITE; PS00237; G PROTEIN RECEP F1 1; 1.
PROSITE; PSS0262; G_PROTEIN_RECEP_F1_2; 1.
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05,
24,
                                                                                                                                                (TrEMBLrel. 19,
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Best Local Similarity 96.8
Matches 30; Conservative
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01-JAN-1998 (TrEMBLrel.
01-JUN-2003 (TrEMBLrel.
                                                                                                       PRELIMINARY;
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                                                                                                                                                  01-DEC-2001
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                                                                                                                             .Q95NC3;
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                                                                                                         Q95NC3
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                                                                 RESULT 2
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AIDS Res. Hum. Retroviruses 0:0-0(1997).
BMBL; ARD11541; AB865741.1; -
GO; GO:0011541; C:integral to membrane; IEA.
GO; GO:00018022; F:rhodopsin-like receptor activity; IEA.
GO; GO:0001894; F:rhodopsin-like receptor protein signalin. .; IEA.
InterPro; IPR000276; GPCR_Rhodpsn.
Pfam; PF00001; 7tm 1; 1.
PRINTS; PR000217; GGCRRHODOPSN.
PROSITE; PS00237; GFCRTBN RECEP_F1_1; 1.
PROSITE; PS00237; G_PROTEIN_RECEP_F1_2; 1.
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Mammalia; Butheria; Primates; Catarrhini; Hominidae; Pan.
NCBI_TaxID=9598;
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352 AA; 40598 MW; 39688FA7004C952F CRC64;
                                                                        40466 MW; 3FFFAC7ABAE1D4FB CRC64;
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Last annotation update)
                                                                                                        ); Pred, No. 4.4e-15; 1; Mismatches 0:
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PROSITE; PS00237; G PROTEIN RECEP_F1_1; 1. PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
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NCBI_TaxID=9534;
[1]
                                                                                                                                                                                                                                                                                                                                                                                                                                                       troglodytes (Chimpanzee).
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96.8%;
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01-MAY-2000 (TrEMBLrel. 13,
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05,
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nes 30; Conservative
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352 AA;
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SEQUENCE FROM N.A.
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01-JAN-1998
                                         Receptor.
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SEQUENCE
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EMBL; AF349683; AAK69685.1;
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GO; GO:0004872; F:receptor activity; IEA.
GO; GO:0001584; F:rhodopsin-like receptor activity; IEA.
GO:0007186; F:G-protein coupled receptor protein signalin. . .; IEA.
InterPro; IRR000276; GPCR_Rhodopsn.
Pfam; PF00001; 7tm 1; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Palacios E., Digilio L., McClure H.M., Chen Z., Marx P.A., Goldsmith M.A., Grant R.M.; "Parallel evolution of CCR5-null phenotypes in humans and in a natural host of simian immunodeficiency virus.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SPECIES-C.torquatus torquatus; STRAIN=RCM411;
Bleer B.B., Kuiken C.L., Tooze Z., Foley B.T., Goeken R.M., Brown C.R.,
St Claire M., Hirsch V.M.,
"Characterization of novel simian immunodeficiency viruses from
redcapped mangabeys from Nigeria (SIVrcmNgM411 and Ng9409).";
Submitted (FBB-2001) to the EMBL/GenBank/DDBJ databases.
EMBL; AF094753; AAG62474.1;
EMBL; AF094773; AAG1194.1;
EMBL; AF084003; AAG62471.1;
                                                                                                                                                                                                                                                                                           Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SPECIES—C. torquatus torquatus; STRAIN=1049, and 997;
Chen Z., Kwon D., Jin Z., Monard S., Telfer P., Jones M., Lu C.,
Agullar R., Ho D.D., Marx P.A.;
"Natural infection of a homozygous delta 24 CCR5 red-capped mangabey
with a R2D-tropic simian immunodeficiency virus.";
J. Exp. Med. 0:0-0(1998).
        Cercocebus torquatus atys (Red-crowned mangabey) (Sooty mangabey). Bukaryota, Metazoca, Chordata, Craniata, Vertebrata, Euteleostomi, Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae, Cercocebus.
                                                                                                                                                                                                                                                                                           ;
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                                                                                                                                                                                                                                                                   Length 352;
                                                                                                                                                                                                                                                                                         Indels
                                                                                                                                                                                                                                        352 AA; 40588 MW; 0A4E4119FAC8EC75 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                    01-MAY-2000 (TrEMBLrel. 13, Created)
01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-UUN-2003 (TrEMBLrel. 24, Last annotation update)
                                                                                                                                                                                                                                                                97.0%; Score 159; DB 6; 96.8%; Pred. No. 4.4e-15;
                                                                                                                                                                                                                                                                                                              1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                        31
                                                                                                                                                                                                                                                                                                                                                                                                   344 AA
                                                                                                                                                                                                                                                                       96.8%; Prea. ...
                                                                                                                                                                                    PRINTS; PR00237; GPCRRHODDPSN.
PROSITE; PS00237; G_PROTEIN_RECEP_F1_1; 1.
PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
                                                                                                                                                                                                                                                                                                                                   1 MDYQVSSPTYDINYYTSEPCQKINVKQIAAR
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Cercocebus torguatus torguatus, and
                                                                                                                                                                                                                                                                                                                                                                                                   PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                SEQUENCE FROM N.A.
SPECIES=C.torquatus atys;
                                                                                                                                                                                                                                                                          Local Similarity 96.8 tes 30; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                           Chemokine receptor CCR5.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           NCBI_TaxID=81944, 9531;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Curr. Biol. 0:0-0(1998)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SEQUENCE FROM N.A.
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                                                                                                                                                                                                                              Receptor.
SEQUENCE
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Q9TQR8
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SEQUENCE FROM N.A.

STRAIN=1208, and 009;

A chear Z., Kwon D., Jin z., Monard S., Telfer P., Jones M., Lu C.,

Adular R., Ho D.D., Marx P.A.;

"Natural infection of a homozygous delta 24 CCR5 red-capped mangabey

"Yatural infection of a homozygous delta 24 CCR5 red-capped mangabey

"Yatural infection of a homozygous delta 24 CCR5 red-capped mangabey

"Yatural infection of a homozygous delta 24 CCR5 red-capped mangabey

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"Yatural infection of a homozygous delta 24 CCR5 red-capped mangabey

"Yatural infection of a homozygous delta 24 AA; "J9578 WW; OCEC0547CZF6DE6 CRC64;
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Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae;
Cercopithecinae; Cercocebus.
GO; GO:0016021; C:integral to membrane; IEA.
GO; GO:0004872; F:receptor activity; IEA.
GO; GO:000184; F:rhodopsin-like receptor activity; IEA.
GO; GO:000186; P:rhodopsin-like receptor protein signalin.
InterPro; IPR000276; GPCR_Rhodpsn.
FFam; PF00001; 7cm 1; 1.
FRINTS; PR00021; 7cm 1; 1.
FRINTS; PR00023; G-PROTEIN RECEP F1_1; 1.
PROSITE; PS500237; G-PROTEIN RECEP F1_2; 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   1; Indels
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                                                                                                                                                                                                                                                                                                                                                       344 AA; 39592 MW; E15F5F601191A4D1 CRC64;
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01-NOV-1998 (TrEMBLrel. 08, Last sequence update)
01-UND-2003 (TrEMBLrel. 24, Last annotation update)
Chemokine receptor CRRS.
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Last annotation update)
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                                                                                                                                                                                                                                                                                                                                                                                                                    ch 93.9%; Score 154; DB 6; 1 Similarity 93.5%; Pred. No. 2.3e-14; 29; Conservative 1; Mismatches 1.
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352 AA

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STRAIN=52;
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SEQUENCE
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                 Q9TV48;
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Q9TV48
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                                                                                                                                                Gaps
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Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hylobatidae, Hylobates.
             Eukaryota, Metazoa, Chordata; Craniata, Vertebrata, Euteleostomi, Mammalia; Butheria; Primates; Catarrhini; Cercopithecidae; Cercopithecus.

NCBI_TaxID=9534;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Submitted (AUG-1999) to the EMBL/GenBank/DDBJ databases.

EMBL; AF177884; AAK43367.1; -...

GO; GO:0016621; C:integral to membrane; IEA.

GO; GO:000485; F:receptor activity; IEA.

GO; GO:0001884; F:rhodopsin-like receptor activity; IEA.

GO; GO:0007186; P:G-protein coupled receptor protein signalin.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 352;
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                                                                                                                                                                                                                                                                                                                                                                                                                            40588 MW; OF1869D9A6668DBB CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          01-DEC-2001 (TrEMBLrel. 19, Created)
01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 154; DB 6;
Pred, No. 2.4e-14;
    (Grivet)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1; Mismatches
                                                                                                                                                                                                                                                                                                                                                                 PROSITE; PS00237; G PROTEIN RECEP_F1_1; 1. PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
      Cercopithecus aethiops (Green monkey)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PRT;
                                                                                                                                                                                                                                                                                                                       Pfam; PF00001; 7tm 1; 1. PRINTS; PR00237; GPCRRHODOPSN.
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93.5%;
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Best Local Similarity 93.5
Matches 29; Conservative
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Best Local Similarity 93.5'
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PROSITE; PS00237; G PR
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                                                                                                                                                                                                                                                                                                                                                                                                                              352 AA;
                                                                                                                                 SEQUENCE FROM N.A.
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SEQUENCE
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RESULT 10

Q9TV48

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Mueller-Trutwin M.C., Corbet S., Hansen J., Diop O.,
Mueller-Trutwin M.C., Corbet S., Hansen J., Diop O.,
Georges-Courbot M.-C., Barre-Sinoussi F., Fomsgaard A.;
Submitted (NOV-1997) to the EMBL/GenBank/DDBJ databases.
EMBL, AR035216; AAD44009.1; --
GO; GO:0004812; F:receptor activity; IEA.
GO; GO:0004815; F:receptor activity; IEA.
GO; GO:0001584; F:rhodopsin-like receptor activity; IEA.
GO; GO:0007186; P:G-protein coupled receptor protein signalin. .; IEA.
PRINTS; PR00237; GFCRRHODOPSN.
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Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi;
Mammalia, Butheria, Primates, Catarrhini, Cercopithecidae,
                                                                                                                                                                                   Eukaryota, Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Butheria; Primates; Catarrhini; Cercopithecidae;
Cercopithecinae; Cercopithecus.
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                                                                                                                                                                                                                                                                                                                                                        MEDLINE=99335215; PubMed=10408730; Muller-Trutwin M.C., Corbet S., Hansen J., Georges-Courbot M.C., Diop O., Rigoulet J., Barre-Sinoussi F., Fomsgaard A.; "Mutations in CKB-coding sequences are not associated with SIV carrier status in African nonluman primates."; AIDS Res. Hum. Retroviruses 15:931-939(1999).
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Pred. No. 2.4e-14;
                                                                                                                                                               Cercopithecus ascanius (black-cheeked white-nosed monkey).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1; Indels
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01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
CC chemokine receptor type 5 (C-C chemokine receptor 5)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       40650 MW; 7906256AA3945266 CRC64;
                                        01-MAY-2000 (TrEMBLrel. 13, Created)
01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
CC chemokine receptor type 5.
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93.5%; Pred. No. 4..
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      PROSITE; PS00237; G PROTEIN RECEP F1 1; 1. PROSITE; PS50262; G PROTEIN RECEP F1 2; 1.
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PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Cercopithecinae; Cercocebus NCBI_TaxID=9532;
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PRELIMINARY;
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Cercopithecus patas.
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                                                                                                            Receptor.
                                                                                                                                        SEQUENCE
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018770
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1D 099
1D 099
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1D 011
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                                                                                                                                                                                                                                                                                                Submitted (AUG-1999) to the EMBL/GenBank/DDBJ databases.

BMBL; AR035115, AAD44008.1; -.

BMBL; AR173896; AAM43381.1; -.

GO; GO:0016021; C:integral to membrane; IEA.

GO; GO:001604; F:rhodopsin-like receptor activity; IEA.

GO; GO:001894; F:rhodopsin-like receptor protein signalin. .; IEA.

InterPro; IPR000276; GPCR_Rhodpsn.

InterPro; IPR00027; GPCR_Rhodpsn.

PRINTS; PR00237; GPCR_RHODPSN.

PROSITE; PS00237; GPCR_RHODPSN.

PROSITE; PS00237; GPCR_RHODPSN.

PROSITE; PS00237; GPCR_RHODPSN.
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Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae,
Cercopithecinae, Cercopithecus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   SPECIES=C.lhoest;
MEDLINE=99335215; PubMed=10408730;
Muller-Trucwin M.C., Corbet S., Hansen J., Georges-Courbot M.C.,
Diop O., Rigoulet J., Barre-Sinoussi F., Fomsgaard A.;
"Mutations in CCRS-coding sequences are not associated with SIV
carrier status in African nonhuman primates.";
AIDS Res. Hum. Retroviruses 15:931-939(1999).
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                                                                                                                                                                             SEQUENCE FROM N.A. Zhang Y.; Zhang Y.; Sequence comparison of the CCRS gene in primates and primate
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          93.9%; Score 154; DB 6; Length 35:
93.5%; Pred. No. 2.4e-14;
"". "". "" Indels
                                                       Mueller-Trutwin M.C., Corbet S., Hansen J., Diop O.,
Georges-Courbot M.-C., Barre-Sinoussi F., Fomsgaard A.,
Submitted (NOV-1997) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              352 AA; 40479 MW; 5E1504A9B87278B2 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-NOV-1999 (TrEMBLrel. 12, Created)
01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Cercopithecus lhoesti (L'Hoest's monkey), and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               1 MDYQVSSPIYDINYYTSEPCOKINVKQIAAR 31
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SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                                 phylogeny."
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      DAT DOTH BY SERVICE STREET STR
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Hull subtypes, co-receptor usage, and CCR5 polymorphism.";
AIDS Res Hum. Retroviruses 0:0-0(1997).
EMBL, AF011538, AAB65738.1;
GO; GO:00016021; C:integral to membrane; IEA.
GO; GO:0004872; F:receptor activity, IEA.
GO; GO:0001584; F:receptor activity, IEA.
GO; GO:0001584; F:rhodopsin-like receptor activity; IEA.
GO; GO:0001584; F:rhodopsin-like receptor protein signalin. .; IEA.
InterPro; IPR000276; GPCR_Rhodops.
Pfam; PR0001; 7tm 1; 1.
PRINTS; PR00237; GPRRHODOPSN.
PROSITE; PS00237; GPRRHODOPSN.
PROSITE; PS00262; GPROTEIN_RECEP_F1.1; 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Pan.
NCBI_TaxID=9598;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Butheria; Primates; Catarrhini; Cercopithecidae;
Cercopithecinae; Cercopithecus.
NCBL_TaxID=27677;
[1]
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                                                                                                                                                               Length 352;
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STRAIN=MacCR5-140a;
Zhang L., Carruthers C.D., He T., Huang Y., Cao Y., Wang Ho D.D.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               1; Indels
                                                                                                          352 AA; 40408 MW; 2D354E5128C779E0 CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  01-JAN-1998 (TrEMBLrel. 05, Created)
01-JAN-1998 (TrEMBLrel. 05, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-MAY-2000 (TrEMBLrel. 13, Created)
01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
CC chemokine receptor type 5.
                                                                                                                                                         93.9%; Score 154; DB 6;
93.5%; Pred. No. 2.4e-14;
vative 1; Mismatches 1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        93.9%; Score 154; DB 6; 93.5%; Pred. No. 2.4e-14;
                                                                                                                                                                                                                                                                    1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                  1 MDYQVSSPTYDIDYYTSEPCQKINVKQIAAR 31
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PRINTS; PR00237; GPCRRHODOPSN.
PROSITE; PS00237; G_PROTEIN_RECEP_FI_1; 1.
PROSITE; PS50262; G_PROTEIN_RECEP_FI_2; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1 MDYQVSSPTYDIDYYTSEPCQKINVKQIAAR
                                                                                                                                                                                                                                                                                                                                                                                                                                                      PRT;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Pan troglodytes (Chimpanzee).
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                                                                                                                                                                                                                29; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                      PRELIMINARY;
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Mueller-Trutwin M.C., Barre-Sinoussi F., Fomsgaard A.;
Georges-Courbot M.-C., Barre-Sinoussi F., Fomsgaard A.;
Submitted (NOV-1997) to the EMBL/GenBank/DDBJ databases.
EMBL, AF035220; AAD44013.1; --
GO; GO:0016021; C:integral to membrane; IEA.
GO; GO:0004822; F:receptor activity; IEA.
GO; GO:0001884; F:rhodopsin-like receptor activity; IEA.
GO; GO:0007186; P:d-protein coupled receptor protein signalin. .; IEA.
Interpro; IPR00027; GPCR_Rhodopsn.
PROSITE; PS00037; GPCRRHODOPSN.
PROSITE; PS50262; G_PROTEIN_RECEP_F1_1; 1.
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GO; GO:0004872; F:receptor activity; IEA.
GO; GO:0001884; F:rhodopsin-like receptor activity; IEA.
GO; GO:0001186; F:G-protein coupled receptor protein signalin. . .; IEA.
InterPro; IPR000276; GPCR_Rhodpsn.
Pfam; PF00001; 7tm_l; 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Zhang Y.-W., Zhang Y.-P.; "Sequence evolution of chemokine receptor CCRS gene in primates."; Submitted (JUN-1998) to the EMBL/GenBank/DDBJ databases.
EMBL; AF075450; AAD19862.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Macaca arctoides (Stump-tailed macaque), and
Macaca assamensis (Assam's macaque) (Assam's monkey).
Eukaryota; Metazoo; Chordata; Craniata; Verrebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae;
Cercopithecinae; Macaca.
                              MEDLINE=99335215; PubMed=10408730; Muller-Trutwin M.C., Corbet S., Hansen J., Georges-Courbot M.C., Diop O., Rigoulet J., Barre-Sinoussi F., Fomsgaard A.; Mutations in CCRS-coding sequences are not associated with SIV carrier status in African nonhuman primates."; AIDS Res. Hum. Retroviruses 15:931-939(1999).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 154; DB 6; Length 35
Pred. No. 2.4e-14;
1; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     40521 MW; SF276C85909FACB2 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                 352 AA; 40747 MW; 3A56E90D3528D94C CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                01-MAY-1999 (TrEMBLrel. 10, Created)
01-MAY-1999 (TrEMBLrel. 10, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
C-C chemokine receptor type 5.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            352 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             EMBL; AF075449; AAD19861.1; -.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                    93.9%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                              93.97
Best Local Similarity 93.55
Matches 29, Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity 93.5
Matches 29; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         352 AA;
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                                                                                                                                                       SEQUENCE FROM N.A.
SEQUENCE FROM N.A
                                                                                                                                                                          STRAIN=04;
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                                                                                                                                                                                                                                                                                                                                                                                                                                   SEQUENCE
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8 3

Search completed: July 29, 2004, 13:29:44 Job time : 36 secs

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GenCore version 5.1.6
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Copyright (c) 1993 - 2004 Compugen Ltd.	using sw model	July 29, 2004, 13:27:36; Search time 18 Seconds (without alignments) 88.911 Million cell updates/sec
Copyright (c) 1	OM protein - protein search, using sw model	July 29, 2004, 13
	M protein -	tun on:

US-09-852-238A-5

164 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR Perfect score: Sequence:

**BLOSUM62** 

Scoring table:

31

389414 seqs, 51625971 residues Gapop 10.0 , Gapext 0.5 Searched:

389414 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

/cgn2\_6/ptodata/2/iaa/5A\_COMB.pep:\*
/cgn2\_6/ptodata/2/iaa/5B\_COMB.pep:\*
/cgn2\_6/ptodata/2/iaa/6A\_COMB.pep:\*
/cgn2\_6/ptodata/2/iaa/6B\_COMB.pep:\*
/cgn2\_6/ptodata/2/iaa/PcTUS\_COMB.pep:\*
/cgn2\_6/ptodata/2/iaa/PcTUS\_COMB.pep:\* Issued Patents AA:\* Database :

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

# SUMMARIES

Description	Segmence 18. Appl	15,	equence 4, 1	17,	equence 6,	equence 13,	Sequence 14, Appl	~	'n	7	equence 1,	52,	52,	2,	Sequence 5, Appli	4	32,	Sequence 32, Appl	32,	'n	9	7	14,	741	19,	19,	19,
ΩI	US-09-087-232A-18	-087-2	US-08-833-752-4	-09-087-2	US-08-833-752-6	19-087-	861-1	3-575-9	US-08-833-752-5	US-09-502-783A-2	US-09-796-202-1	-09 - 045		-08-466	US-09-517-605-5	US-08-861-105-4	-08-88-	-09-098-244-3	-09-375-3	-60-	0	-08-724-9	US-09-131-827A-14	US-09-621-976-7413	US-08-663-566A-19	US-08-023-610-19	US-08-288-065A-19
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Score	164	164	164	164	164	164	164	164	164	164	164	S	159	2	4	123	Н	$\vdash$	113	113	105	10	81.5	52	51	51	51
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Sequence 19, Appl	Sequence 19, Appl	4, A	13		Sequence 50, Appl	6	Sequence 9, Appli	θ,	. 4	4		51,	Sequence 50, Appl	51	7	ς,	Sequence 20, Appl
US-08-362-240A-19	PCT-US95-10245-19	US-09-242-435-4	US-09-131-827A-13	US-09-117-257-50	US-09-489-352-50	US-09-502-783A-9	US-08-466-343D-9	US-08-461-244-3	US-08-450-393A-4	US-08-446-669-4	US-09-045-583-50	US-09-045-583-51	US-09-534-185-50	US-09-534-185-51	US-08-833-752-7	US-09-131-827A-2	US-09-131-827A-20
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705	705	1130	83	170	170	329	344	347	360	360	360	360	360	360	360	360	360
31.1	31.1	31.1	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9	29.9
51	21	21	49	49	4	4	49	49	49	49	49	49	49	49	49	49	40
28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

## ALIGNMENTS

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Sequence 18, Application US/09087232A
Patent No. 6153431
GENERAL INFORMATION:
TITLE OF INVENTION: HUMAN IMMUNOBERCIENCY VIRUS CO-RECEPTOR
TITLE OF INVENTION: VARIANTS ASSOCIATED WITH RESISTANCE TO VIRUS INFECTION.
NUMBER OF SEQUENCES:
ADDRESSEE: Baker & Botts, L.L.P. attn. Lisa Kole
STREET: 30 Rockefeller Plaza
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                0
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                                                                                                                                                                                                                                                                                                       ZIP: 1012
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: EUR PC Compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/087,232A
FILING DATE: 28 MAY 1998
CLASSIFICATION: 435
PRIOR APPLICATION: A75
PRIOR DATE: 30 MAY 1997
APPLICATION NUMBER: 60/048,057
FILING DATE: 30 MAY 1997
ATTORNEY/AGENT INFORMATION:
NAME: 30 MAY 1997
REFERENCE/DOCKET NUMBER: 35,225
REFERENCE/DOCKET NUMBER: AP 31115
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               100.0%; Score 164; DB 3;
100.0%; Pred. No. 1e-17;
cive 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TELECOMMUNICATION INFORMATION:
TELEPHONE: (212) 408-2628
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TELEFAX: (212) 765-2519
INFORMATION FOR SEQ ID NO: 1
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             LENGIH: 87 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           31; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  amino acid
3Y: linear
                                                                                                                                                                                                                                             CITY: New York
STATE: New York
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
Matches 31; Conserv
US-09-087-232A-18
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TELEPHONE:
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US-09-087-232A-17
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                                                                                                                                                                                                                                                                                                                                                            US-08-833-752-4
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                                     Sequence 15, Application US/09087232A

Patent No. 6153431

GENERAL INFORMATION:

APPLICANT: Quillent et al.

TITLE OF INVENTION: "HUMAN IMMUNOBERICIENCY VIRUS CO-RECEPTOR

TITLE OF INVENTION: VARIANTS ASSOCIATED WITH RESISTANCE TO VIRUS INFECTION.

NUMBER OF SEQUENCES: (23.

CORRESPONDENCE ADDRESS:

ADDRESSER: Baker & Botts, L.L.P. attn. Lisa Kole

STREET: 30 Rockefeller Plaza
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ö
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APPLICANT: PARENTIER, MARC
APPLICANT: PARENTIER, MARC
APPLICANT: LIBERT, GILBERT
APPLICANT: LIBERT, GILBERT
TITLE OF INVENTION: ACTIVE AND INACTIVE CC-CHEMOKINES RECEPTOR
TITLE OF INVENTION: AND NUCLEIC ACID MOLECULES ENCODING SAID RECEPTOR
NUMBER OF SEQUENCES: 17
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                            SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO) CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/087,232A
FILING DATE: 28 MAY 1998
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 164; DB 3;
Pred. No. 1.2e-17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADDRESSEE: Knobbe, Martens, Olson & Bear
STREET: 620 Newport Center Drive 16th Floor
CITY: Newport Beach
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      100.0%; Scott No. 100.0%; Pred. No. 100.0%; Pred. No. 100.0%; Pred. No. 100.0%; Nismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  REFERENCE/DOCKET NUMBER: AP 31115
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION DATA:
FILING DATE: 30 MAY 1997
ATTONEY/ABOTT INFORMATION:
NAME: KOLE, LISA B.
REGISTRATION NUMBER: 35,225
                                                                                                                                                                                                                                                                                                                    ZIP: 10112
COMPUTER REALABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequence 4, Application US/08833752
Patent No. 6448375
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TELECOMMUNICATION INFORMATION TELEPHONE: (212) 408-2628
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity 100.0%
Matches 31, Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (212) 765-2519
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                INFORMATION FOR SEQ ID NO: SEQUENCE CHARACTERISTICS: LENGTH: 100 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ; TOPOLOGY: linear; MOLECULE TYPE: protein US-09-087-232A-15
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      GENERAL INFORMATION:
                                                                                                                                                                                                                                                       CITY: New York STATE: New York
                                                                                                                                                                                                                                                                                                      USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      COUNTRY: U
RESULT 2
US-09-087-232A-15
                                                                                                                                                                                                                                                                                               COUNTRY:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 3
US-08-833-752-4
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Sequence 17, Application US/09087232A
Patent No. 615443+
GENERAL INFORMATION:
APPLICANT: Quillent et al.
TITLE OF INVENTION: HUMAN IMMUNOBERICIENCY VIRUS CO-RECEPTOR
TITLE OF INVENTION: VARIANTS ASSOCIATED WITH RESISTANCE TO VIRUS INFECTION.
NUMBER OF SEQUENCES: 23
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                100.0%; Score 164; DB 4; Length 184; 100.0%; Pred. No. 2.5e-17; tive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CURKENENDEMICK ADDRESS:
ADDRESSEE: Baker & Botts, L.L.P. attn. Lisa Kole
STREET: 30 Rockefeller Plaza
CITY: New York
STATE: New York
COUNTRY: USA
ZIP: 10112
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPUTER: DEM PC compatible
COMPUTER: DATA:
APPLICATION NUMBER: 06/048,057
RILING DATE: 28 MAY 1997
ATTORNEY/AGENT INFORMATION:
NAME: FOLLS: DISA B.
REFERENCE/DOCKET NUMBER: AP 31115
TELECOMMUTCATION NUMBER: AP 31115
TELECOMMUTCATION NUMBER: AP 31115
COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC Compatible
COMPUTER: IBM PC Compatible
COMPUTER: IBM PC COMPATIBLE
COMPATION TYPE: PLOS/MS-DOS
SOFTWARE: PATENTIN Release #1.0, Version #1.25 (EPO)
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/833,752
FILING DATE: 9-APR-1997
CLASSIFICATION: 536
ATTORNEY/AGENT INFORMATION:
NAME: Altura, Daniel E
REGISTRATION NUMBER: 34,115
REGISTRATION NUMBER:
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
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INFORMATION FOR SEQ ID NO: 1
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                             LENGTH: 184 amino acids
TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    31; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MOLECULE TYPE: protein
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 amino acid
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Matches 31; Conserv
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TELEPHONE:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 7
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                                                                                           0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ·.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  APPLICANT: SARSON, MICHEL
APPLICANT: PARMENTIER, MARC
APPLICANT: PASSART, GILBERT
APPLICANT: LIBERT, FREDERICK
TITLE OF INVENTION: ACTIVE AND INACTIVE CC-CHEMOKINES RECEPTOR
TITLE OF INVENTION: AND NUCLEIC ACID MOLECULES ENCODING SAID RECEPTOR
TORRESPONDENCE ADDRESS:
ADDRESSEE: Knobbe, Martens, Olson & Bear
                                                                                       Gaps
                                                                                       ·
0
100.0%; Score 164; DB 3; Length 215; 100.0%; Pred. No. 3e-17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            100.0%; Score 164; DB 4; Length 215;
llarity 100.0%; Pred. No. 3e-17;
Conservative 0; Mismatches 0; Indels
                                                                                   Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   SOFTWARE: Patentin Release #1.0, Version #1.25 (EPO) CURRENT APPLICATION DATA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            E: Baker & Botts, L.L.P. attn. Lisa Kole
30 Rockefeller Plaza
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADDRESSEE: Knobbe, Martens, Olson & Bear
STREET: 620 Newport Center Drive 16th Floor
CITY: Newport Beach
                                                                                                                                                  1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                             1 MDYQVSSPIYDINYYTSEPCOKINVKOIAAR 31
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                                                                               0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  APPLICATION NUMBER: US/08/833,752
FILING DATE: 9-APR-1997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                COMPUTER READABLE FORM:
MEDLUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    US-09-087-232A-13
Sequence 13, Application US/09087232A
Patent No. 6153431
GENERAL INFORMATION:
TAPLICAN: Quillent et al.
TITLE OF INVENTION: HUMAN IMMUNODI
TITLE OF INVENTION: VARIANTS ASSOC
                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 6, Application US/08833752
Patent No. 6448375
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    NAME: Altman, Daniel E
REGISTRATION NUMBER: 34,115
REFERENCE/DOCKET NUMBER:
INFORMATION FOR SEQ ID NO: 6:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 FILING DATE: 9-APR-1997
CLASSIFICATION: 536
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      : 215 amino acids amino acid
                                   Local Similarity 100. es 31; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ; MOLECULE TYPE: protein US-08-833-752-6
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     U.S.A.
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STATE: New York
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Simi
Matches 31;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TOPOLOGY:
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Query Match
                                                                       Matches
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APPLICANT: DISTANCE DENG, HONGKUI
APPLICANT: BLAMEIER, WILFRIED
APPLICANT: LANDAU NATHANIEL R.
APPLICANT: LIU, RONG
TITLE OF INVENTION: G-COUPLED RECEPTORS ASSOCIATED WITH
TITLE OF INVENTION: MACROPHAGE-TROPHIC HIV, AND DIAGNOSTIC AND THERAPEUTIC
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS: 7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ·;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Indels
            MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC Compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30 (BPO)
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/087,232A
FILING DATE: 28 MAY 1998
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADDRESSEE: David A. Jackson, Esq.
STREET: 411 Hackensack Ave, Continental Plaza, 4th
STREET: Floor
STREET: Roor
STREET: May Jersey
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PatentIn Release #1.0, Version #1.30
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100.0%; Score 164; DB 3;
Best Local Similarity 100.0%; Pred. No. 5.4e-17;
Matches 31; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                   PILING DATE: 28 MAY 1998
CLASSIFICATION: 435
PRIOR APPLICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 60/048,057
FILING DATE: 30 MAY 1997
ATTORNEY/AGENT INFORMATION:
NAME: KOLE, LISA B.
REFERENCE/DOCKET NUMBER: 35,225
REFERENCE/DOCKET NUMBER: AP 3111:
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/861,105
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  COMPUTER READBBLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IEM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 14, Application US/08861105
Patent No. 6258272
GENERAL INFORMATION:
APPLICANT: LITTMAN, DAN R.
                                                                                                                                                                                                                                                                                                                                                                                                                                           TELEPAX: (212) 765-2519
INFORMATION FOR SEQ ID NO: 13:
SEQUENCE CHARACTERISTICS:
LENGTH: 352 amino acids
TYPE: amino acid
TOPOLOGY: linear
                                                                                                                                                                                                                                                                                                                                                                                                                            (212) 408-2628
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CLASSIFICATION: 436
PRIOR APPLICATION DATA:
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ; MOLECULE TYPE: protein US-09-087-232A-13
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PRIOR APPLICATION DATA:
APPLICATION NUMBER: U
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Pred. No. 5.4e-17;

100.08;

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31; Conservative
Best Local Similarity
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US-09-502-783A-2
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      STATE:
                       Matches
                                                                     à
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Sequence 2, Application US/08575967A
Patent No. (Settle of the organization US/08575967A
Patent No. (Settle of the organization US/08575967A

Tarke of invention: Chemokine Receptor Materials and Methods NUMBER OF SEQUENCES: 16
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADDRESSEE: Marshall, O'Toole, Gerstein, Murray & Borun STREET: 6300 Sears Tower, 233 S. Wacker Drive
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Indels
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COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM FC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/575,967A
FILING DATE:
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: NO. 6265184and, Greta E.
REGISTRATION NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 35,302
REGISTRATION NUMBER: 35,302
REGISTRATION NUMBER: 35,302
REGISTRATION NUMBER: 35,302
REGISTRATION NUMBER: 1662
TELEPHONE: 206-485-1900
TELEPHONE: 206-485-1900
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OTHER INFORMATION: /= "88C amino acid sequence"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
100.0%; Score 164; DB 3;
Best Local Similarity 100.0%; Pred. No. 5.4e-17;
Matches 31; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1 MDYQVSSPIYDINYYTSEPCOKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                         FILING DATE: 13-APR-1994
ATTONNEY/ABENT INDORMATION:
NAME: Jackson Esq., David A.
REGISTRATION NUMBER: 26,742
REFERENCE/DOCKET NUMBER: 1049-1-004 NI
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION 18FORMATION:
         US 08/227,319
                                                                                                                                                                                TELEFAX: 201-343-1684
INFORMATION FOR SEQ ID NO: 14:
SEQUENCE CHARACTERISTICS:
LENGTH: 352 amino acids
TYPE: amino acid
STRANDEDNESS: single
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TYPE: amino acids
TOPOLOGY: 1:-CIPE OFFER
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                                                                                                                                                                                                                                                                                                                                                                                                             Homo sapiens
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                                                                                                                                                                                                                                                                                                                                            MOLECULE TYPE: protein HYPOTHETICAL: NO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Chicago
: Illinois
RY: USA
                                                                                                                                                                                                                                                                                                                         linear
                                                                                                                                                                                                                                                                                                                                                                                      ORIGINAL SOURCE:
CORGANISM: Home
US-08-861-105-14
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CITY: Chi
STATE: Il
COUNTRY:
                                                                                                                                                                                                                                                                                                                         TOPOLOGY:
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DB 3; Length 352;

100.0%; Score 164;

Query Match

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TITLE OF INVENTION: POLYMOLGOCIDES Encoding Human G-Protein Chemokine Receptor (CCRETILE OF INVENTION: HDGNR10
FILE REFERENCE: 1488.1150006
CURRENT APPLICATION NUMBER: U5/09/502,783A
CURRENT FILING DATE: 2001-08-23
PRIOR FILING DATE: 1955-06-06
NUMBER OF SEQ ID NOS: 9
SOFTWARE: Patentin version 3.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ö
                                                                                                                                                                                                                        APPLICANT: SAMSON, MICHEL
APPLICANT: PARRENTIER, MARC
APPLICANT: PARRENTIER, MARC
APPLICANT: GILBERT
APPLICANT: GILBERT
TITLE OF INVENTION: ACTIVE AND INACTIVE CC-CHEMOKINES RECEPTOR
TITLE OF INVENTION: AND NUCLEIC ACID MOLECULES ENCODING SAID RECEPTOR
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
Gaps
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Indels
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100.0%; Pred. No. 5.4e-17;
iive 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                     ADDRESSEE: Knobbe, Martens, Olson & Bear
STREET: 620 Newport Center Drive 16th Floor
CITY: Newport Beach
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
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                                                          1 MDYQVSSPIYDINYTSEPCQKINVKQIAAR 31
 0; Mismatches
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                                                                                                                                                  US-08-833-752-5
; Sequence 5, Application US/08833752
Patent No. 6448375
; GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    INFORMATION FOR SEQ ID NO: 5: SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         : 352 amino acids
amino acid
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Best Local Similarity 100.
Matches 31; Conservative
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MOLECULE TYPE: protein US-08-833-752-5
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1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                REFERENCE/DOCKET NUMBER: MNI-044
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Best Local Similarity
Matches 30; Conserva
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Patent No. 6287805
GENERAL INFORMATION:
APPLICANT: Graham, Gerard J. et al.
TITLE OF INVENTION: No. 6287805el Molecules of the G Protein-Coupled NUMBER OF SEQUENCES: 56
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD, LLP
STREET: 28 State Street
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
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0
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Batent No. 6548636

GENERAL INFORMATION:
APPLICANT: Disaic, Tatjana
APPLICANT: Olson, William
TITLE OF INVENTION: SULFATED CCRS PEPTIDES FOR HIV-1 INFECTION
FILE REPERENCE: 2048/61010/JPW/SHS
CURRENT APPLICATION NUMBER: US/09/796,202
CURRENT PILING DATE: 2001-02-28
NUMBER: PALENTING DATE: 2001-02-28
SOFTWARE: PALENTIN VERSION 3.0
SEQ ID NO 1
                                                                                                                           Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 352;
                                                                                                                                                                Indels
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COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM FC Compatible
OPPRATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
                                                                                                                   100.0%; Score 164; DB 4;
100.0%; Pred. No. 5.4e-17;
iive 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         100.0%; Score 164; DB 4;
100.0%; Pred. No. 5.4e-17;
tive 0; Mismatches 0;
                                                                                                                                                                                              1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                       1 MDYQVSSPIYDINYYTSEPCOKINVKQIAAR 31
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      NAME: Mandragouras, Amy E. REGISTRATION NUMBER: 36,207
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         FILING DATE:
ATTORNEY/AGENT INFORMATION:
                                                                                                                                      Best Local Similarity 100.0
Matches 31; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              31; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CITY: Boston
STATE: Massachusetts
                                      TYPE: PRT
ORGANISM: Homo sapiens
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity
Matches 31; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          USA
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; ORGANISM: human
US-09-796-202-1
                                                                             US-09-502-783A-2
; SEQ ID NO 2
; LENGTH: 352
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  US-09-045-583-52
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        LENGTH: 352
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        COUNTRY:
                                                                                                                     Query Match
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RESULT 13
US-09-534-185-52
; Sequence 52, Application US/09534185
; Patent No. 6403767;
GENERAL INFORMATION:
; APPLICANT: Graham, Gerard J. et al.
TITLE OF INVENTION: No. 6403767el Molecules of the G Protein-Coupled Title OF INVENTION: Metafor
                                                                                                                                                                                                                                                                                                               Gaps
                                                                                                                                                                                                                                                               Score 159; DB 3; Length 352; Pred. No. 3.1e-16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC Compatible
COMPUTER: IBM PC Compatible
COMPUTER: IBM PC Compatible
COMPUTER: Batentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/534,185
FILING DATE: 24-Mar-2000
CLASSIFICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION NUMBER: 09/045,583
FILING DATE: cunknown-
ATTORNEY/AGENT INFORMATION:
NAMME: MANDER: AMY E.
REGISTRATION NUMBER: 36,207
REFERENCE/DOCKET NUMBER: MI-044
TELECOMMUNICATION INFORMATION:
TELEBHONE: (617)227-7400
                                                                                                                                                                                                                                                                                                          0; Indels
                                                                                                                                                                                                                                      Query Match 97.0%; Score 159; DB Best Local Similarity 96.8%; Pred. No. 3.1e
                                                                                                                                                                                                                                                                                                                                                                            1 MDYQVSSPIYDIDYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                 1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADDRESSEE: LAHIVE & COCKFIELD, LLP
STREET: 28 State Street
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ; MOLECULE TYPE: peptide
; FRAGMENT TYPE: internal
SEQUENCE DESCRIPTION: SEQ ID NO: 52:
US-09-534_185-52
                        TELEFAX: (617)742-4212
INFORMATION FOR SEQ ID NO: 52: SEQUENCE CHARACTERISTICS: LENGTH: 352 amino acids
TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              LENGTH: 352 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (617) 742-4214
TELECOMMUNICATION INFORMATION TELEPHONE: (617) 227-7400
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 STATE: Massachusetts
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 INFORMATION FOR SEQ ID NO: 52:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   COMPUTER READABLE FORM:
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                                                                                                                                                TOPOLOGY: linear
MOLECULE TYPE: peptide
; FRAGMENT TYPE: internal
US-09-045-583-52
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CITY: Boston
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ZIP: 02109
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Gaps

0

97.0%; Score 159; DB 4; Length 352; 96.8%; Pred. No. 3.1e-16; Live 1; Mismatches 0; Indels

1 MDYQVSSPIYDIDYYTSEPCQKINVKQIAAR 31

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Sequence 5, Application US/09517605

Sequence 5, Application US/09517605

Sequence 5, Application US/09517605

Barent No. 6391567

GENERAL INFORMATION:
APPLICANT: Littman, Dan R.
APPLICANT: Woon, Douglas S.
APPLICANT: Van Kooyk, Yvette
APPLICANT: Geijtenbeck, Theo
TITLE OF INVENTION: CELLS
FILE REFERENCE: 1049-1-017

CURRENT APPLICATION NUMBER: US/09/517,605

CURRENT FILING DATE: 2000-03-02

NUMBER OF SEQ ID NOS: 17

SOGTWARKE: Patentin Ver. 2.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                            96.3%; Score 158; DB 3; Length 352; 96.8%; Pred. No. 4.5e-16; tive 0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1 MDYQVSSPIYDINYYTSEPCQKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1 MDYQVSSPIYDINYYTSEPCPKINVKQIAAR 31
                                                                                                                                                                                                                                                                                                                                                                             : 352 amino acids
amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity 96.8
Matches 30; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                             MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                   TOPOLOGY:
                                                                                                                                                                                                                                                                                                                                                                                                                         US-08-466-343D-2
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US-09-517-605-5
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TYPE: PRT ORGANISM: Homo sapiens

SEQ ID NO 5 LENGTH: 352

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                           Query Match

87.8%; Score 144; DB 4; Length 352;
Best Local Similarity 90.3%; Pred. No. 6e-14;
Matches 28; Conservative 1; Mismatches 2; Indels
                                                                                                                            1 MDYQVSSPIYDINYYTSEPCOKINVKQIAAR 31
US-09-517-605-5
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Search completed: July 29, 2004, 13:30:40 Job time : 19 secs